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[Continued on next page]

#### (54) Title: REGULATION OF EMBRYONIC TRANSCRIPTION IN PLANTS

-950	ACTCA	TAAAAACTAG	TAGATTGGTT	GGTTGGTTTC	CATGTACCAG
				ATDIOFW >	
-900	AAGGCTTACC	CTATTAGTTG			TACTCAATTC
-850	CTAGTTGTGT	AAATGTATGT	ATATGTAATG	CGTATAAAAC	GTAGTACTTA
-800	AATGACTAGG	AGTGGTTCTT	GAGACCGATG	AGAGATGGGA	GCAGAACTAA
-750	AGATGATGAC	ATAATTAAGA	ACGAATTIGA	AAGGCTCTTA	GGTTTGAATC
-700	CTATTCGAGA	ATGTTTTTGT	CARAGATAGT	GGCGATTTTG	AACCAAAGAA
-650	AACATTTAAA	AAATCAGTAT	CCGGTTACGT	TCATGCAAAT	AGAAAGTGGT
-600	CTAGGATCTG	ATTGTAATTT	TAGACTTAAA	GAGTCTCTTA	AGATTCAATC
-550	CTGGCTGTGT	ACAAAACTAC	AAATAATATA	TTTTAGACTA	TTTGGCCTTA
-500	ACTAAACTTC	CACTCATTAT	TTACTGAGGT	TAGAGAATAG	ACTTGCGAAT
-450	AAACACATTC	CCGAGAAATA	CTCATGATCC	CATAATTAGT	CAGAGGGTAT
-400	GCCAATCAGA	TCTAAGAACA	CACATTCCCT	CARATTTTAR	TGCACATGTA
-350	ATCATAGTTT	AGCACAATTC	AAAAATAATG	TAGTATTAAA	GACAGAAATT
-300	TGTAGACTTT	TTTTTGGCGT	TAAAGGAAGA	CTAAGTTTAT	ACGTACATT?
-250	TATTTTAAGT	GGAAAACCGA	AATTTTCCAT	CGAAATATAT	GAATTTAGTA
-200	TATATATTTC	TGCAATGTAC	TATTTTGCTA	TTTTGGCAAC	TTTCAGTGGA
-150	CTACTACTTT	ATTACAATGT	GTATGGATGC	ATGAGTTTGA	GTATACACAT
-100	GTCTAAATGC	ATGCTTTGCA	AAACGTAACG	GACCACAAAA	GAGGATCCAT
-50	GCAAATACAT	CTCATAGCTT	CCTCCATTAT		AAACAGAGCA
10	ATGACGTCCG	TTARCGTTAR		Atproky	

GATTCAATC
TIGGCCTTA
CTTGCGAAT
AGAGGGTAT
GCACATGTA
ACAGAAATT
CGTACATTT

1 ATGACGTCCG TTAACGTTAA GCTCCTT

(57) Abstract: Nucleic acid constructs are provided comprising transcriptional regulatory regions homologous to plant FAE1 promoters. In some embodiments, these constructs may be used in transgenic cells or plants to promote expression of foreign and endogenous genes in developing seeds, for example to affect seed lipid metabolism, protein or carbohydrate composition and accumulation, or seed development.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

#### REGULATION OF EMBRYONIC TRANSCRIPTION IN PLANTS

#### FIELD OF THE INVENTION

The invention is in the field of nucleic acid sequences capable of regulating transcription. particularly sequences that may promote transcription during embryogenesis in plants.

#### **BACKGROUND OF THE INVENTION**

Most of the information about seed-specific gene expression comes from studies of genes encoding seed storage proteins like napin, a major protein in the seeds of *Brassica napus*, or conglycinin of soybean. Upstream DNA sequences directing strong embryo-specific expression of these storage proteins have been used successfully in transgenic plants to manipulate seed lipid composition and accumulation (Voelker et al., 1996). However. expression of storage protein genes begins fairly late in embryogenesis. Thus, promoters of seed storage protein genes may not be ideal for all seed-specific applications. For example, storage oil accumulation commences significantly before the highest level of expression of either napin (Stalberg et al., 1996) or conglycinin (Chen et al., 1988) is achieved. It is, therefore of interest to identify other promoters which may modulate expression of genes in developing plant embryos.

A variety of transcriptional regulatory regions that may be active during plant embryogenesis are known, as disclosed for example in: U.S. Patent No. 5,792,922 issued 11 August 1998 to Moloney; U.S. Patent No. 5,623,067 issued 22 April 1997 to Vandekerckhove et al.: International Patent Publication WO9845461 published 15 October 1998. There remains a need for alternative transcriptional regulatory regions.

FATTY ACID ELONGATION1 (FAE1) genes encode condensing enzymes involved in plant very long chain fatty acid biosynthesis. The FAE1 condensing enzyme is thought to be localized in the endoplasmic reticulum where it catalyzes the sequential elongation of C18 fatty acyl chains to C22 in length (Kunst et al., 1992). FAE1 genes have been cloned and described recently by James et al. (1995), International Patent Publication WO 96/13582.

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#### SUMMARY OF THE INVENTION

In one aspect, the invention provides transcriptional regulatory regions derived from *FAE1* genes. The transcriptional regulatory regions of the invention may be useful in

promoting early seed-specific transcription of heterologous sequences to which they are operably linked. The transcriptional regulatory regions of the invention may be used in a wide variety of plants, including *Brassica sp.*, *Arabidopsis* and other plant species. DNA constructs comprising the transcriptional regulatory sequences of the invention may be active during fatty acid or lipid biosynthesis in the plant embryo. Certain embodiments of the constructs of the invention may be used in transgenic plants to promote expression of heterologous sequences in developing seeds. In various embodiments, the constructs of the invention may be used to mediate gene expression that affects seed lipid metabolism, or seed protein composition or seed carbohydrate composition, or seed development. In alternative embodiments, the transcriptional regulatory regions of the invention may also be useful for the production of modified seeds containing novel recombinant proteins which have pharmaceutical, industrial or nutritional value.

#### BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 shows a 934 bp DNA sequence comprising the *Arabidopsis thaliana FAE1* transcription regulatory sequence.

**Figure 2** shows a 1588 bp DNA sequence comprising the *Brassica napus FAE1* transcription regulatory sequence.

**Figure 3** shows a 1069 bp DNA sequence comprising the *Lunaria annua FAE1* transcription regulatory sequence.

Figure 4 shows an alignment of the *Arabidopsis thaliana* (A.t.), *Lunaria annua* (L.a.) and *Brassica napus* (B.n.) transcription regulatory sequences. Asterisks below the sequences indicate identical nucleotides in each of the three sequences. A number of putative cis-acting sequence motifs are identified in the A. thaliana sequence: an EM1 ABA box at -44bp to -36bp having the sequence ACATCTCAT, for which the published consensus sequence is ACGTGTCAT (Rowley, D.L. and Herman, E.M. (1997), Biochimica et Biophysica Acta 1345:1-4); an A-300 box at -51bp to -46bp having the sequence TGCAAT, for which the published consensus sequence is TG(T/A/C)AAA(G/T) (Morton et al. (1994) in Seed Development and Germination (Kigel, J. and Gallili, G., eds.) pp. 103-138, Marcel Dekker. New York); G-box 1 at -105 to -100 bp having the sequence CACATG, for which is the consensus sequence is CACCTG, and G-box 2 at -164 to -159 bp having the sequence CAACTT, for which the consensus sequence is CAACTG (Kawogoe, Y. and Murai, N. (1992) Plant J. 2:927-936; CE1 element at -226 to -218 bp having the sequence

TTCCATCGA, for which the consensus sequence is TGCCACCGG, and a CE3 element at -381bp to -369 bp having the sequence ACACATTCCCTC, for which the consensus sequence is ACGCGTGTCCTC (Shen et al., (1996) Plant Cell 8:1107-1119). Not highlighted is a putative RY repeat motif at -53bp to -47bp having the sequence CATGCAA, for which the consensus sequence is CATGCAT (Dickinson et al. (1988) Nucleic Acid Res. 16:371; Lelievre et al. (1992) Plant Physiol. 98:387-391). Also shown, as Con. 4, is a consensus sequence, wherein R=G or A, Y=T/U or C, M=A or C, K=G or T/U, S=G or C, W=A or T/U, B=G or C or T/U, D=A or G or T/U, H=A or C or T/U, V=A or G or C and N=A or G or C or T/U.

Figure 5 shows an alignment of the Arabidopsis thaliana (A.t.) and Lunaria annua (L.a.) transcription regulatory sequences. Asterisks below the sequences indicate identical nucleotides in each of the two sequences. The base at position -400 in the A.t. sequence is highlighted. The alignment of sequences in both Figure 4 and Figure 5 was accomplished using the CLUSTALW program (version 1.74) for multiple sequence alignments, using a gap open penalty of 15, a gap extension penalty of 6.66 and an IUB DNA weight matrix. Also shown, as Con. 5, is a consensus sequence, wherein R=G or A, Y=T/U or C, M=A or C, K=G or T/U, S=G or C, W=A or T/U, B=G or C or T/U, D=A or G or T/U, H=A or C or T/U, V=A or G or C and N=A or G or C or T/U.

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Figure 6 includes two bar graphs illustrating hydroxy fatty acid content of A) FAE120 FAH12 and B) napin-FAH12 transgenic seeds, expressed as percentage of total seed fatty acids.

Figure 7 shows an alignment of the *Brassica napus* (B.n.) and *Lunaria annua* (L.a.) FEA1 transcription regulatory sequences. Asterisks below the sequences indicate identical nucleotides in each of the two sequences.

Figure 8 shows an alignment of the *Brassica napus* (B.n.) and *Arabidopsis thaliana* (A.t.) FEA1 transcription regulatory sequences. Asterisks below the sequences indicate identical nucleotides in each of the two sequences.

#### DETAILED DESCRIPTION OF THE INVENTION

The recombinant nucleic acid molecules of the invention may comprise a heterologous promoter sequence operably linked to a nucleic acid sequence, wherein the promoter sequence comprises a transcriptional regulatory region capable of mediating seed-specific expression in *Arabidopsis*. The transcriptional regulatory region may be obtainable from a

plant FAE1 gene. Alterntively, The transcriptional regulatory region may hybridize under stringent conditions to a 5' region of the plant FAE1 gene. In further alternative embodiments, The transcriptional regulatory region may be at least 70% identical when optimally aligned to the 5' region of the plant FAE1 gene.

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In alternative embodiments, the invention provides isolated nucleic acids comprising the transcriptional regulatory regions of the invention. By isolated, it is meant that the isolated substance has been substantially separated or purified away from other biological components with which it would otherwise be associated, for example *in vivo*. The term 'isolated' therefore includes substances purified by standard purification methods, as well as substances prepared by recombinant expression in a host, as well as chemically synthesized substances.

In the context of the present invention, "transcriptional regulatory region" means a nucleotide sequence capable of mediating or modulating transcription of a nucleotide sequence of interest, when the transcriptional regulatory region is operably linked to the sequence of interest. Conversely, a transcriptional regulatory region and a sequence of interest are "operably linked" when the sequences are functionally connected so as to permit transcription of the sequence of interest to be mediated or modulated by the transcriptional regulatory region. In some embodiments, to be operably linked, a transcriptional regulatory region may be located on the same strand as the sequence of interest. The transcriptional regulatory region may in some embodiments be located 5' of the sequence of interest. In such embodiments, the transcriptional regulatory region may be directly 5' of the sequence of interest or there may be intervening sequences between these regions. The operable linkage of the transcriptional regulatory region and the sequence of interest may require appropriate molecules (such as transcriptional activator proteins) to be bound to the transcriptional regulatory region, the invention therefore encompasses embodiments in which such molecules are provided, either *in vitro* or *in vivo*.

The term "recombinant" means that something has been recombined, so that when made in reference to a nucleic acid molecule the term refers to a molecule that is comprised of nucleic acid sequences that are joined together by means of molecular biological techniques. The term "recombinant" when made in reference to a protein or a polypeptide refers to a protein molecule which is expressed using a recombinant nucleic acid molecule. The term "heterologous" when made in reference to a nucleic acid sequence refers to a nucleotide sequence which is ligated to, or is manipulated to become ligated to, a nucleic acid sequence

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to which it is not ligated in nature, or to which it is ligated at a different location in nature. The term "heterologous" therefore indicates that the nucleic acid molecule has been manipulated using genetic engineering, i.e. by human intervention.

Sequences may be derived or obtainable from plant FAE1 genes by deduction and synthesis based upon the wild-type FAE1 gene sequences. Derived sequences may be identified in different organisms, for example by isolation using as probes the nucleic acid sequences of the invention. Alternative transcriptional regulatory regions may be derived through mutagenesis or substitution of wild-type sequences, such as the sequence disclosed herein. Derived nucleic acids of the invention may be obtained by chemical synthesis, isolation, or cloning from genomic DNAs using techniques known in the art, such as the Polymerase Chain Reaction (PCR). Consensus sequences, such as those illustrated in Figures 4 and 5 are alternative embodiments of the nucleic acids of the invention, derived from the disclose wild-type FAE1 gene sequences. Nucleic acids of the present invention may be used to design alternative primers (probes) suitable for use as PCR primers to amplify particular regions of an FAE1 gene. Such PCR primers may for example comprise a sequence of 15-20 consecutive nucleotides of the sequences of the invention. To enhance amplification specificity, primers of 20-30 nucleotides in length may also be used. Methods and conditions for PCR amplification are described in Innis et al. (1990); Sambrook et al. (1989); and Ausubel et al. (1995). As used herein, the term "probe" when made in reference to an oligonucleotide refers to an oligonucleotide which is capable of hybridizing to another oligonucleotide of interest. A probe may be single-stranded or double-stranded. Probes are. for example, useful in the detection, identification, amplification and isolation of particular gene sequences. Oligonucleotide probes may be labelled with a "reporter molecule." so that the probe is detectable using a detection system, such as enzymatic, fluorescent, radioactive or luminescent detection systems.

Derived nucleic acids of the invention may also be identified by hybridization, such as Southern or Northern analysis. Southern analysis is a method by which the presence of DNA sequences in a target nucleic acid mixture are identified by hybridization to a labeled probe, comprising an oligonucleotide or DNA fragment of a nucleic acid of the invention. Probes for Southern analysis may for example be at least 15 nucleotides in length. Southern analysis typically involves electrophoretic separation of DNA digests on agarose gels, denaturation of the DNA after electrophoretic separation, and transfer of the DNA to nitrocellulose, nylon, or another suitable membrane support for analysis with a radiolabeled, biotinylated, or enzyme-

labeled probe as described in Sambrook *et al.* (1989). Similarly, Northern analysis may be used to identify RNAs that hybridize to a known probe such as an oligonucleotide, DNA fragment. cDNA or fragment thereof, or RNA fragment of a nucleic acid of the invention or a known *FAE1* sequence. The probe may be labeled with a radioisotope such as <sup>32</sup>P, by biotinylation or with an enzyme. The RNA to be analyzed may be electrophoretically separated on an agarose or polyacrylamide gel, transferred to nitrocellulose, nylon, or other suitable membrane, and hybridized with the probe, using standard techniques well known in the art such as described in Sambrook *et al.* (1989).

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In alternative embodiments, a transcriptional regulatory region of the invention may be at least 70% identical when optimally aligned to the 5' region of a plant FAE1 gene, such as the Arabidopsis FAE1 gene. In alternative embodiments, the degree of identity may be between 50% and 100%, such as 60%, 80%, 90%, 95% or 99%. When a position in the compared sequence is occupied by the same nucleotide or amino acid. following optimal alignment of the sequences, the molecules are considered to have identity at that position. The degree of identity between sequences is a function of the number of matching positions shared by the sequences. In terms of percentage, identity is the sum of identical positions, divided by the total length over which the sequences are aligned, multiplied by 100.

Various aspects of the present invention encompass nucleic acid or amino acid sequences that are homologous to other sequences. As the term is used herein, an amino acid or nucleic acid sequence is "homologous" to another sequence if the two sequences are substantially identical and the functional activity of the sequences is conserved (for example, both sequences function as or encode a FAE1 enzyme: as used herein, the term 'homologous' does not infer evolutionary relatedness). Nucleic acid sequences may also be homologous if they encode substantially identical amino acid sequences, even if the nucleic acid sequences are not themselves substantially identical, a circumstance that may for example arise as a result of the degeneracy of the genetic code.

Two amino acid or nucleic acid sequences are considered substantially identical if, when optimally aligned (with gaps permitted), they share at least about 50% sequence similarity or identity, or if the sequences share defined functional motifs. In alternative embodiments, sequence similarity in optimally aligned substantially identical sequences may be at least 60%, 70%, 80%, 90% or 95%. As used herein, a given percentage of homology between sequences denotes the degree of sequence identity in optimally aligned sequences.

Optimal alignment of sequences for comparisons of similarity may be automated using a variety of algorithms, such as the local homology algorithm of Smith and Waterman (1981) Adv. Appl. Math 2: 482, the homology alignment algorithm of Needleman and Wunsch (1970) J. Mol. Biol. 48:443, the search for similarity method of Pearson and Lipman (1988) Proc. 5 Natl. Acad. Sci. USA 85: 2444, and the computerized implementations of these algorithms (such as GAP, BESTFIT, FASTA and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, Madison, WI, U.S.A.). Sequence similarity may also be determined using the BLAST algorithm, described in Altschul et al. (1990), J. Mol. Biol. 215:403-10 (using the published default settings). Software and instructions for performing 10 BLAST analysis may be available through the National Center for Biotechnology Information in the United States (including the programs BLASTP, BLASTN, BLASTN, TBLASTN and TBLASTX that may be available through the internet at http://www.ncbi.nlm.nih.gov/). The BLAST algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence that either match or satisfy some positivevalued threshold score T when aligned with a word of the same length in a database 15 (reference) sequence. T is referred to as the neighborhood word score threshold. Initial neighborhood word hits act as seeds for initiating searches to find longer HSPs. The word hits are extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Extension of the word hits in each direction is halted when the following parameters are met: the cumulative alignment score falls off by the quantity X from 20 its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T and X determine the sensitivity and speed of the alignment. The BLAST program may use as defaults a word length (W) of 25 11, the BLOSUM62 scoring matrix (Henikoff and Henikoff (1992) Proc. Natl. Acad. Sci. USA 89: 10915-10919), a gap existence cost of 11, a per residue gap cost of 1, a lambda ratio of 0.85, alignments (B) of 50, expectation (E) of 10, M=5, N=4, and a comparison of both strands. One measure of the statistical similarity between two sequences using the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the 30 probability by which a match between two nucleotide or amino acid sequences would occur by chance. In alternative embodiments of the invention, nucleotide or amino acid sequences are considered substantially identical if the smallest sum probability in a comparison of the test sequences is less than about 1, preferably less than about 0.1, more preferably less than

about 0.01, and most preferably less than about 0.001. In the PSI-BLAST implementation of the BLAST algorithm, an expect value for inclusion in PSI-BLAST iteration may be 0.001 (Altschul et al. (1997), Nucleic Acids Res. 25:3389-3402). Searching parameters may be varied to obtain potentially homologous sequences from database searches.

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An alternative indication that two nucleic acid sequences are substantially identical is that the two sequences hybridize to each other under moderately stringent, or preferably stringent, conditions. Hybridization to filter-bound sequences under moderately stringent conditions may, for example, be performed in 0.5 M NaHPO<sub>4</sub>, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65EC, and washing in 0.2 x SSC/0.1% SDS at 42EC (see Ausubel, et al. (eds), 1989, Current Protocols in Molecular Biology, Vol. 1, Green Publishing Associates. Inc., and John Wiley & Sons, Inc., New York, at p. 2.10.3). Alternatively, hybridization to filter-bound sequences under stringent conditions may, for example, be performed in 0.5 M NaHPO<sub>4</sub>, 7% SDS, 1 mM EDTA at 65EC, and washing in 0.1 x SSC/0.1% SDS at 68EC (see Ausubel, et al. (eds), 1989, supra). Hybridization conditions may be modified in accordance with known methods depending on the sequence of interest (see Tijssen, 1993, Luboratory Techniques in Biochemistry and Molecular Biology -- Hybridization with Nucleic Acid Probes, Part I, Chapter 2 "Overview of principles of hybridization and the strategy of nucleic acid probe assays", Elsevier, New York). Generally, stringent conditions are selected to be about 5EC lower than the thermal melting point for the specific sequence at a defined ionic strength and pH.

A FAE1 promoter is any naturally occurring transcriptional regulatory region that mediates or modulates the expression of a plant FAE1 condensing enzyme. Plant FAE1 condensing enzymes are proteins that are homologous to known FAE1 condensing enzymes, such as those cloned and described in International Patent Publication WO 96/13582.

Heterologous DNA sequences may for example be introduced into a host cell by transformation. Such heterologous molecules may include sequences derived from the host cell species, which have been isolated and reintroduced into cells of the host species. Heterologous nucleic acid sequences may become integrated into a host cell genome, either as a result of the original transformation of the host cells, or as the result of subsequent recombination events. Transformation techniques that may be employed include plant cell membrane disruption by electroporation, microinjection and polyethylene glycol based transformation (such as are disclosed in Paszkowski et al. EMBO J. 3:2717 (1984); Fromm et al., Proc. Natl. Acad. Sci. USA 82:5824 (1985); Rogers et al., Methods Enzymol. 118:627

(1986); and in U.S. Patent Nos. 4,684,611; 4,801,540; 4,743,548 and 5,231,019), biolistic transformation such as DNA particle bombardment (for example as disclosed in Klein, et al., Nature 327: 70 (1987); Gordon-Kamm, et al. "The Plant Cell" 2:603 (1990); and in U.S. Patent Nos. 4,945,050; 5,015,580; 5,149,655 and 5,466,587); Agrobacterium-mediated transformation methods (such as those disclosed in Horsch et al. Science 233: 496 (1984); Fraley et al., Proc. Nat'l Acad. Sci. USA 80:4803 (1983); and U.S. Patent Nos. 4,940,838 and 5,464,763).

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Standard methods are available for the preparation of constructs for use in identifying and characterizing transcriptional regulatory regions useful in various embodiments of the invention. General molecular techniques may for example be performed by procedures generally described by Ausubel FM, Brent R, Kingston RE, Moore DD, Seidman JG, Smith JA, Stuhl K. (1995) Current Protocols in Molecular Biology, Vols 1, 2 and 3. Alternative equivalent methods or variations thereof may be used in accordance with the general knowledge of those skilled in this art and the functional requirements of the present invention.

In some aspects of the invention, transformed plant cells may be cultured to regenerate whole plants having a transformed genotype and displaying a desired phenotype, as for example modified by the expression of a heterologous protein mediated by a transcriptional regulatory region of the invention. A variety of plant culture techniques may be used to regenerate whole plants, such as are described in Gamborg and Phillips, "Plant Cell, Tissue and Organ Culture, Fundamental Methods", Springer Berlin, 1995); Evans et al. "Protoplasts Isolation and Culture", Handbook of Plant Cell Culture, Macmillian Publishing Company, New York, 1983; or Binding, "Regeneration of Plants, Plant Protoplasts", CRC Press, Boca Raton. 1985: or in Klee et al., Ann. Rev. of Plant Phys. 38:467 (1987). A cell. tissue. organ, or organism into which has been introduced a foreign nucleic acid, is considered "transformed", "transfected", or "transgenic". A transgenic or transformed cell or organism also includes progeny of the cell or organism and progeny produced from a breeding program employing a transgenic plant as a parent in a cross and exhibiting an altered phenotype resulting from the presence of a recombinant nucleic acid construct. A transgenic plant is therefore a plant that has been transformed with a heterologous nucleic acid, or the progeny of such a plant that includes the transgene. The invention provides vectors, such as vectors for transforming plants or plant cells. The term "vector" in reference to nucleic acid molecule generally refers to a molecule that may be used to transfer a nucleic acid segment(s) from one cell to another. One

of skill will recognize that after the nucleic acid is stably incorporated in transgenic plants and

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confirmed to be operable, it can be introduced into other plants by sexual crossing. Any of a number of standard breeding techniques may be used, depending upon the species to be crossed.

In various embodiments, the invention comprises plants transformed with the nucleic acids of the invention. In some embodiments, such plants will exhibit altered fatty acid content in one or more tissues. These aspects of the invention relate to all higher plants, including monocots and dicots, such as species from the genera Fragaria. Lotus, Medicago, Onobrychis, Triforium, Trigonelia, Wgna, Citrus, Linum. Geranium, Manihot. Caucus, Arabidopsis, Brassica, Raphanus, Sinapis, Atropa, Capsicum, Hyoscyamus. Lycopersicon. Nicotiana, Solanum, Petunia, Digitalis, Majorana, Cichorium, Helianthus, Lactuca, Bromus, Asparagus, Antirrhinum, Heterocatlis, Nemesia, Pelargonium, Panicum, Penniserum. Ranunculus, Senecio, Salpiglossis, Cucarnis, Browallia, Glycine, Lolium, Zea, Triticum, Sorghum. and Datura. Such plants may include maize, wheat, rice, barley, soybean, beans. rapeseed, canola, alfalfa, flax, sunflower, cotton, clover, lettuce, tomato cucurbits, potato carrot, radish. pea lentils, cabbage, broccoli, brussel sprouts, peppers, apple, pear, peach. apricot, carnations and roses. More specifically, in alternative embodiments, plants for which the invention may be used in modifying fatty acid content include oil crops of the Cruciferae family: canola, rapeseed (Brassica spp.), crambe (Crambe spp.), honesty (Lunaria spp.) lesquerella (Lesquerela spp.), and others; the Composirae family: sunflower (Helianthus spp.), safflower (Carthamus spp.), niger (Guizotia spp.) and others; the Palmae family: palm (Elaeis spp.), coconut (Cocos spp.) and others; the Leguminosae family: peanut (Arachis spp.), soybean (Glycine spp.) and others: and plants of other families such as maize (Zea spp.), cotton (Gossvpiun sp.), jojoba (Simonasia sp.), flax (Linum sp.), sesame (Sesamum spp.), castor bean (Ricinus spp.), olive (Olea spp.), poppy (Papaver spp.), spurge (Euphorbia, spp.), meadowfoam (Limnanthes spp.), mustard (Sinapis spp.) and cuphea (Cuphea spp.).

Nucleic acids of the invention may also be used as a plant breeding tool, as molecular markers to aid in plant breeding programs. Such techniques would include using the gene itself as a molecular probe or using the DNA sequence to design PCR primers to use PCR based screening techniques in plant breeding programs.

Deletion or insertion constructs may be useful for domain mapping to determine the functional domains or motifs of a transcriptional regulatory region derived from a FAE1 gene. An aspect of the invention is the construction and testing of such constructs, as described below for the 5' deletion construct of the A. thaliana FAE1 5' region. One aspect of the

invention comprises transcriptional regulatory regions that are derived from functionally important regions of a FAE1 promoter. As outlined above, the functionally important regions of a FAE1 promoter may be determined through routine assays. Alternatively, randomly selected portions of a a FAE1 promoter may be selected for use in routine assays to determine whether the selected region is capable of functioning as a transcriptional regulatory region in the context of the present invention. In various embodiments, regions of the Arabidopsis thaliana, Brassica napus or Lunaria annua promoters may be used. For example, the following motifs in the A.t. FAE1 promoter may be used alone or in combination in novel transcriptional regulatory regions (see Figure 4): the CE-like elements (CE1 and CE3), the RY repeat motif, the G-boxes (G-box1 and G-box2), the A-300 box, the EM1 ABA box. or the CTATTTTG element. Constructs of the invention comprising such motifs, deletions or insertions may be assayed for activity as transcriptional regulatory regions of the invention by testing for strong seed-specific activity providing expression of a sequence of interest (such as a reporter sequence) before the torpedo stage and persisting throughout embryo development. in accordance with standard testing methods that may be adapted from the methods disclosed herein.

Alternative embodiments of the transcriptional regulatory regions of the invention may be identified using information available through NCBI databases at http://www.ncbi.nih.gov.

In various embodiments, transcriptional regulatory regions derived from plant FAE1 genes are shown to be capable of directing expression of desired genes at an early stage of development in a seed-specific manner in disparate plant species. In particular embodiments, the transcriptional regulatory regions of the invention may be used in a wide variety of dicotyledonous plants for modification of the seed phenotype. For example, new seed phenotypes may include:

- 25 (1) altered seed fatty acid composition or seed oil composition and accumulation
  - (2) altered seed protein or carbohydrate composition or accumulation
  - (3) enhanced production of desirable endogenous seed products

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- (4) suppression of production of undesirable gene products using antisense. co-suppression or ribozyme technologies
- 30 (5) production of novel recombinant proteins for pharmaceutical, industrial or nutritional purposes

Isolation of a seed-specific promoter from A. thaliana

Using the sequence information of the *A. thaliana* genome sequencing project. synthetic oligonucletide primers were designed to amplify the *FAE1* 5' untranslated region, to isolate it by PCR. As shown in Figure 1, the upstream primer 5'-CTAGTAGATTGGTTGGTTGGTTTCC-3' (AtproFW) in combination with the downstream primer 5'-TGCTCTGTTTGTGTCGGAAAATAATGG-3' (AtproRV) were used, and resulted in the synthesis of a fragment of the correct size (934 bp). The amplified product was subcloned in the *Hin*cII site of the plasmid pT7T3-18U (Pharmacia) to produce plasmid pT7T3-18U/proFAE900, followed by complete sequence determination of both strands to verify the fragment identity. A BLAST search of the *A. thaliana* Database identified a single BAC clone T4L20 (GenBank ATF10M6) 125,179 bp long, which contains the complete *FAE1* gene.

#### Functional analysis of the FAE1 5' upstream region

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5' upstream fragments of the FAEI gene were shown to confer seed-specific and temporally regulated gene expression in plants. A translational fusion was made between the FAEI 5' region and the coding region of the reporter gene  $\beta$ -glucuronidase (GUS). The chimeric gene (pFAE900-GUS or pFAE400-GUS) was transferred into *Arabidopsis* and tobacco and GUS activity was monitored in various tissue of transgenic plants.

Construction of the vectors pFAE900-GUS and pFAE400-GUS, and transformation of *Arabidopsis* and tobacco, was as follows. The insert was cleaved out of pT7T3-18U vector with *Hin*dIII and *Xba*I and directionally subcloned into the corresponding sites of the binary Ti plasmid pBI101 (Clontech), which contains a promoterless GUS gene (Jefferson et al. 1987), to obtain the vector pFAE900-GUS. Another construct, pFAE400-GUS, containing only 393 bp of the 5' *FAE1* region directly upstream of the ATG initiation codon fused to the GUS coding sequence was also generated. For that, the pT7T3-18U/proFAE900 vector was digested with *BglII* and *PstI*, the sticky ends were filled in using T4 DNA polymerase, followed by re-ligation to obtain pT7T3-18U/proFAE400. The 393 bp 5' *FAE1* upstream fragment was then excised with *Hin*dIII and *Xba*I and cloned into the binary vector pBI101 to obtain the plasmid pFAE400-GUS. The pFAE400-GUS and pFAE900-GUS fusion constructs in pBI101 were introduced into *Agrobacterium tumefaciens* strain GV3101 (Koncz and Schell, 1986) by heat-shock and selected for resistance to kanamycin (50 µg/ml). *A. thaliana* (L.) Heynh. ecotype Columbia was transformed with the pFAE400-GUS and pFAE900-GUS constructs using floral dip method (Clough and Bent, 1998). Screening for transformed seed

was done on 50μg/mL kanamycin as described previously (Katavic et al., 1994). Approximately 100 transgenic lines were generated for each construct.

For transformation of tobacco, *A. tumefaciens* harbouring the pFAE900-GUS construct was co-cultivated with leaf pieces of *Nicotiana tabacum* SR1 and transformants were selected with kanamycin (100µg/mL) on solid medium (Lee and Douglas, 1996).

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Histochemical localization of GUS activity and analysis of transgenic plants was as follows. Tissue sections were placed in 100 mM NaPO<sub>4</sub> (pH7) and 1 mM spermidine for 15 min, then incubated at  $37^{\circ}$  C in 0.5 K<sub>3</sub>[Fe(CN)<sub>6</sub>], 0.01 % Triton X-100, 1mM EDTA, 10 mM  $\beta$ -mercaptoethanol, 5-bromo-4-chloro-3-indolyl- $\beta$ -D-glucuronide in 100 mM NaPO<sub>4</sub> (pH7), until a blue color appeared (after approximately 1 hr). Following incubation with the substrate, chlorophyll was removed from the sections using a graded ethanol series.

Using this assay, five independent transgenic *Arabidopsis* lines were examined for the embryo-specific expression of the GUS gene. In addition, leaf, stem and siliques were histochemically stained for β-glucuronidase activity. The results indicate that the reporter gene fused to the transcriptional regulatory region of the invention is not expressed in vegetative tissues, whereas it is highly expressed in developing seeds (embryos). Both the 934 bp and the 393 bp transcriptional regulatory regions derived from the *A.t. FAE1* gene caused the appearance of GUS activity by the torpedo stage embryo (6 days after flowering). GUS activity in all five lines persisted throughout subsequent embryo development.

Leaves, stems, pods and seeds of three regenerated tobacco lines transformed with the pFAE900-GUS construct were also assayed for β-glucuronidase activity. The results obtained indicate that the 934 bp *FAE1* promoter fragment contains sufficient information to direct seed-specific expression of a reporter gene in transgenic tobacco. Thus the transcriptional regulatory regions of the invention may be used for seed-specific expression of foreign genes in transgenic plants.

The *in vivo* activity of a *FAE1* promoter of the invention was compared to the activity of the napin promoter by expressing the castor bean hydroxylase gene *FAH12* (Broun and Somerville, 1997) behind either the *FAE1*-promoter (a transcriptional regulatory region of approximately 1 kb) or the napin promoter in an Arabidopsis *fad2/fae1* double mutant. This mutant accumulates as a proportion of fatty acids about 85% of the 18:1 acyl group, which is the substrate for the hydroxylase. The levels of hydroxylated fatty acids accumulating in a large number of independent transgenic lines were used to estimate the relative strength of

each promoter. As shown in Figure 6, the two populations of transgenic plants accumulated levels of hydroxylated fatty acids, ranging from 0.2% to about 11-12% of total fatty acids, with the levels being on average slightly higher in *FAE1-FAH12* lines. Similarly, the best *FAE1-FAH12* plant accumulated just over 12% of hydroxylated fatty acids (w/w of total FAs), whereas the best napin-FAH12 plant produced 10.8% of hydroxylated fatty acids (w/w of total FAs). These results indicate that the FAE1 promoter is highly active in transgenic *Arabidopsis* and that its *in vivo* activity may be superior to napin in *Arabidopsis* seeds.

Sequence elements or motifs that confer both tissue specificity and developmental regulation of transcription reside within 393 bp of the AUG translation initiation codon in the A.t. FAE1 gene. The seed-specific expression conferred by the transcriptional regulatory regions of the invention is independent of the native terminator of the FAE1 gene 3' end. For example, in the exemplified constructs disclosed herein, a terminator derived from the Agrobacterium nopaline synthase gene was used.

#### Lunaria annua and Brassica napus FAE1 5' regulatory regions

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Two sequences originating from B. napus and L. annua were isolated and characterized to demonstrate that regulatory regions conferring seed-specific transcription early in embryo development can also be found upstream of other plant FAE1 genes. Sequences were cloned using the technique of polymerase chain reaction (PCR) walking on uncloned plant genomic DNA (Devic et al., 1997). Approximately 5 µg of genomic DNA from 1 g of fresh tissue was used for the construction of 5 different libraries by digesting DNA with a series of enzymes that produce blunt end fragments to which special adaptors are ligated. The adaptor molecules consist of a long upper strand, which contains successive sequences common to the adaptor primers, AP1 and AP2, annealed at its 3' end to a shorter strand lacking the AP1 sequence. However, this short strand posseses an amine group at its 3' end to prevent filling in by the DNA polymerses during the first PCR amplification step and generation of the AP1 binding site. This suppression PCR effect prevents exponential amplification of molecules containing the adaptor at each end, and the adaptor primer binding sites are only produced when a strand complementary to the upper strand of the adaptor is synthesized by extension from a gene specific primer. The first PCR reaction is performed using an adaptor primer AP1 and a gene specific primer. An aliquot of the first PCR product is used a template in a second PCR amplification using the nested gene specific primer and AP2.

In order to isolate the regulatory regions upstream of the B. napus FAE1 coding sequence, genomic DNA was prepared from developing leaves and digested with 5 blunt-end

cutting restriction enzymes (*Dral. EcoRV, Hpal, Pvull and Scal*) to generate a series of DNA libraries. After ligation of adapter molecules, individual libraries were used as templates in a two step PCR. In the first PCR amplification using the AP1 primer 5'-GGATCCTAATACGACTCACTATAGGGC-3' and the *FAE1* gene specific primer 5'-AAAGAGTGGAGCGATGGTTATGAGG-3' (Bnwalk1), multiple DNA fragments were amplified from all five library templates. After a second round of PCR, using the AP2 primer 5'-CTATAGGGCTCGAGCGGC-3' and the nested *FAE1* specific primer 5'-CGGAAAGAAGCAAAGGTTGAAAAGG-3' (Bnwalk2), the longest single fragment of 1.6 kb was obtained from the *Hpal* library template. This fragment was inserted into the pCR2.1 plasmid (Invitrogen) and sequenced. The sequence is shown in Figure 2.

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For the PCR walking experiment to isolate the *L. annua* 5' regulatory region, in adition to the standard AP1 and AP2 primers, the following *FAE1* specific primers were used: 5'-GATCGTTTGTGGTAAGACGAGAGC-3' (Lawalk1) and 5'-GTCAGTGGGAAGAAACAGAGGTTG-3' (Lawalk2). In the first PCR reaction, the *Dral*, *EcoRV*, *PvuII*, *Sca1* and *Ssp1* library templates were used. In a second PCR amplification the longest single fragment 1.1 kb in length was synthesized using the *EcoRV* library template. This fragment was inserted into the *Hinc11* site of the pT7T3-18U vector (Promega), sequenced on both strands and analyzed (Figure 3).

Using the sequence data obtained for the 5' regulatory regions generated by PCR walking, specific primers were generated for the amplification of the *L. annua* and *B. napus FAE1* promoter fragments. For the PCR-amplification of *B. napus* promoter fragment the upstream primer was 5'-CTGACTTCACCAAAGAAACAACTCG-3' (BnproFW) in combination with the downstream primer 5'-CGGAATTCCGTTTTTTTTTTTAGGCG-3' (BnproRV). The synthesized fragment was ligated into the *Sma1* site of pGEM-7Zf (Promega), then excised with *Xbal/BamHI* and cloned into the equivalent sites of the pBI101 binary vector (Clontech). *L. annua* 5' regulatory region was amplified using the 5'-CAGCTTAACCGGTAAAATTGGCC-3' (LaproFW) upstream primer together with the 5'-TGTTCAGTTTTGTGTCGGAGAGG-3' (LaproRV) downstream primer and inserted in the *Hincl1* site of pT7T3-18U (Promega) plasmid. In order to clone the *L. annua* promoter fragment into the pBI101 binary vector, an *XbaI* site was added by subcloning the *Pstl/KpnI* fragment released from the pT7T3-18U vector into pBluescript II KS+ (Stratagene). The fragment was then excised and cloned in the *XbaI* site of the pBI101 vector.

The resulting vectors pBnFAE1-GUS and pLaFAE1-GUS in pBI101 were then introduced into *A. tumefaciens* strain GV3101 by heat-shock, and used to transform *Arabidopsis* as described above. Transformants were selected on agar-solidified medium containing kanamycin (50 µg/ml). More than 100 transformants were generated for each construct. The activity of the *L. annua* and *B. napus FAE1* promoters was determined by GUS expression assays on the developing seeds and also on non-reproductive plant tissues as controls. Consistent seed-specific GUS expression was obtained for both promoter constructs in independent transgenic lines. In contrast, there was no detectable GUS activity in leaf, stem and silique samples.

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#### WHAT IS CLAIMED IS:

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A recombinant nucleic acid molecule comprising a heterologous promoter sequence
 operably linked to a nucleic acid sequence, wherein the promoter sequence comprises a transcriptional regulatory region capable of mediating seed-specific expression in Arabidopsis wherein the transcriptional regulatory region:

- (a) is obtainable from a 5' region of a plant FAE1 gene; or
- (b) hybridizes under stringent conditions to the 5' region of the plant FAE1 gene; or
- is at least 70% identical when optimally aligned to the 5' region of the plant *FAE1* gene.
- 2. The recombinant nucleic acid of claim 1 wherein the 5' region of the plant FAE1 gene comprises (5' to 3'):

	AGA	TCTAAGAACA	CACATTCCCT	CAAATTTTAA	TGCACATGTA
	ATCATAGTTT	AGCACAATTC	AAAAATAATG	TAGTATTAAA	GACAGAAATT
	TGTAGACTTT	TTTTTGGCGT	TAAAGGAAGA	CTAAGTTTAT	ACGTACATTT
	TATTTTAAGT	GGAAAACCGA	AATTTTCCAT	CGAAATATAT	GAATTTAGTA
20	TATATATTTC	TGCAATGTAC	TATTTTGCTA	TTTTGGCAAC	TTTCAGTGGA
	CTACTACTTT	ATTACAATGT	GTATGGATGC	ATGAGTTTGA	GTATACACAT
	GTCTAAATGC	ATGCTTTGCA	AAACGTAACG	GACCACAAAA	GAGGATCCAT
	GCAAATACAT	CTCATAGCTT CCT	CCATTAT TTTCC	GACAC AAACAGAG	GCA.

The recombinant nucleic acid of claim 1 wherein the 5' region of the plant FAE1 gene comprises (5' to 3'):

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AAGGCTTACC CTATTAGTTG AAAGTTGAAA CTTTGTTCCC TACTCAATTC
CTAGTTGTGT AAATGTATGT ATATGTAATG CGTATAAAAC GTAGTACTTA
AATGACTAGG AGTGGTTCTT GAGACCGATG AGAGATGGGA GCAGAACTAA

30 AGATGATGAC ATAATTAAGA ACGAATTTGA AAGGCTCTTA GGTTTGAATC
CTATTCGAGA ATGTTTTTGT CAAAGATAGT GGCGATTTTG AACCAAAGAA
AACATTTAAA AAATCAGTAT CCGGTTACGT TCATGCAAAT AGAAAGTGGT
CTAGGATCTG ATTGTAATTT TAGACTTAAA GAGTCTCTTA AGATTCAATC
CTGGCTGTGT ACAAAACTAC AAATAATATA TTTTAGACTA TTTGGCCTTA

35 ACTAAACTTC CACTCATTAT TTACTGAGGT TAGAGAATAG ACTTGCGAAT
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AAACACATTC CCGAGAAATA CTCATGATCC CATAATTAGT CAGAGGGTAT
GCCAATCAGA TCTAAGAACA CACATTCCCT CAAATTTTAA TGCACATGTA
ATCATAGTTT AGCACAATTC AAAAATAATG TAGTATTAAA GACAGAAATT
TGTAGACTTT TTTTTGGCGT TAAAGGAAGA CTAAGTTTAT ACGTACATTT
TATTTTAAGT GGAAAACCGA AATTTTCCAT CGAAATATAT GAATTTAGTA
TATATATTTC TGCAATGTAC TATTTGCTA TTTTGGCAAC TTTCAGTGGA
CTACTACTTT ATTACAATGT GTATGGATGC ATGAGTTTGA GTATACACAT
GTCTAAATGC ATGCTTTGCA AAACGTAACG GACCACAAAA GAGGATCCAT
GCAAATACAT CTCATAGCTT CCTCCATTAT TTTCCGACAC AAACAGAGCA.

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4. The recombinant nucleic acid of claim 1 wherein the 5' region of the plant FAE1 gene comprises (5' to 3'):

CTGACTTC ACCAAAGAAA CAACTCGAGT CGTTATCCAT 15 TCTCTACATC GCAACCCGGC CCAAACCGGT TTACCTCGTT GAGTACTCAT GCTACCTTCC ACCAACGCAT TGTAGATCAA GTATCTCCAA GGTCATGGAT ATCTTTTATC AAGTAAGAAA AGCTGATCCT TCTCGGAACG GCACGTGCGA TGACTCGTCG TGGCTTGACT TCTTGAGGAA GATTCAAGAA CGTTCAGGTC TAGGCGATGA AACTCACGGG CCCGAGGGGC TGCTTCAGGT CCCTCCCCGG 20 AAGACTTTTG CGGCGGCGCG TGAAGAGACG GAGCAAGTTA TCATTGGTGC GCTAGAAAAT CTATTCAAGA ACACCAACGT TAACCCTAAA GATATAGGTA TACTTGTGGT GAACTCAAGC ATGTTTAATC CAACTCCATC GCTCTCCGCG ATGGTCGTTA ACACTTTCAA GCTCCGAAGC AACGTAAGAA GCTTTAACCT TGGTGGCATG GGTTGTAGTG CCGGCGTTAT AGCCATTGAT CTAGCAAAGG 25 ACTTGTTGCA TGTCCATAAA AATACGTATG CTCTTGTGGT GAGCACAGAG AACATCACTT ATAACATTTA CGCTGGTGAT AATAGGTCCA TGATGGTTTC AAATTGCTTG TTCCGTGTTG GTGGGGCCGC TATTTTGCTC TCCAACAAGC CTGGAGATCG TAGACGGTCC AAGTACGAGC TAGTTCACAC GGTTCGAACG CATACCGGAG CTGACGACAA GTCTTTTCGT TGCGTGCAAC AAGGAGACGA 30 TGAGAACGGC AAAATCGGAG TGAGTTTGTC CAAGGACATA ACCGATGTTG CTGGTCGAAC GGTTAAGAAA AACATAGCAA CGTTGGGTCC GTTGATTCTT CCGTTAAGCG AGAAACTTCT TTTTTTCGTT ACCTTCATGG GCAAGAAACT TTTCAAAGAT AAAATCAAAC ATTACTACGT CCCGGATTTC AAACTTGCTA TTGACCATTT TTGTATACAT GCCGGAGGCA GAGCCGTGAT TGATGTGCTA 35 GAGAAGAACC TAGCCCTAGC ACCGATCGAT GTAGAGGCAT CAAGATCAAC GTTACATAGA TTTGGAAACA CTTCATCTAG CTCAATATGG TATGAGTTGG

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The recombinant nucleic acid of claim 1 wherein the 5' region of the plant FAE1 gene comprises (5' to 3'):

CAGCTTAAC CGGTAAAATT GGCCTGTACA TATATTTACC ACTGAGTAAA GACATCAGTT AATGATTTGT TGTTACTCAA TTGGGCTAAG TGTATTATTA TATGTGTTGT ATATAATAAA 15 GGTAGAACGT AAATTTACTA AGAATGTGTT TTTCCAATGT GATTGCTCTT TGGCCTCTTA GGTTTGAATC CTACTCGAGA AGACTAATTT TAATTTACTG GCAAAAATAG AAATCAATTT ATAAGTGTTT AAACAAATCG ATGGTATAAC TGATTAGTGA TCACTCTTAG GTTTTGATCC AACTCGAGTA TTGAGTATTG AACGCTTTTT TTAAATAAAA TCTTGATTTT TAAATTGGTT TTTTGAGTAA 20 AAAAGTTCTT AATATTTTCT CTTTGTTTTA ATGGGTTTGT TTTGCATTTT ATAAGCTTAA TTTTTCTAAT TTAATATTTT ATCTATCATC GTCCGTAAAG TTTTATTTGG CACAAACTTG TTTTACTTTT CTACCTTATA ATTTGGGAAC TGGTTGAGTC AAAGCGTACC GGACAAATAT GTTTTATATT CTTATTTAAG AATTAACACT CATCTCATAA TTAGTCAGAG GCTAGGGAGA TTCAGCCAAT 25 CAATGCTAAC AACAAAATTC TCTTAATGAT CTAACGATGC TATTTAATAT TCGGATCAGT ATTCTTAAAT AAGAATATAA AACTAATTCA ATAGTTACAG ATAAAAACTT ATATAGACTT TTTTATTTGG AATATAAAAG TATCAATATA TTATAGACAA TATTTATAAC GTTAAAAATA CAATATTTAT ATTTTTTATA TATTTATTTC AAATTGAAAA GCATTACTTC TATCGAAATG AATTTTAGTA 30 TATTAATTAA TATTTTTTTA ATCGGACTAC TTTCCTATTT TGGCACCTTT CATCTGACTA CTAATTTATT TCAATGTGTA TGCATGCATG AGCATGAGTA ATACACATGT CTATATAAAT GCATGTAAAA CGTAACGGAC CACAAAAGTG GATCCATACA AATACATCTC ATCGCACCCT CTCCGACACA AAACTGAACA.

6. The recombinant nucleic acid of claim 1 wherein the promoter sequence is selected from the group consisting of *Arabidopsis thaliana*, *Lunaria annua* and *Brassica napus FAE1* promoter sequences.

- 5 7. The recombinant nucleic acid of any one of claims 1 through 6, wherein the transcriptional regulatory region is at least 70% identical when optimally aligned to the 5' region of the plant FAE1 gene..
- 8. The recombinant nucleic acid of claim 1 wherein the transcriptional regulatory region comprises (5' to 3'):

	AGA	TCTAAGAACA	CACATTCCCT	CAAATTTTAA	TGCACATGTA
	ATCATAGTTT	AGCACAATTC	AAAAATAATG	TAGTATTAAA	GACAGAAATT
	TGTAGACTTT	TTTTTGGCGT	TAAAGGAAGA	CTAAGTTTAT	ACGTACATTT
	TATTTTAAGT	GGAAAACCGA	AATTTTCCAT	CGAAATATAT	GAATTTAGTA
15	TATATATTTC	TGCAATGTAC	TATTTTGCTA	TTTTGGCAAC	TTTCAGTGGA
	CTACTACTTT	ATTACAATGT	GTATGGATGC	ATGAGTTTGA	GTATACACAT
	GTCTAAATGC	ATGCTTTGCA	AAACGTAACG	GACCACAAAA	GAGGATCCAT
	GCAAATACAT	CTCATAGCTT CCT	CCATTAT TTTCC	GACAC AAACAGAG	CA.

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9. The recombinant nucleic acid of claim 1 wherein the transcriptional regulatory region comprises (5' to 3'):

AAGGCTTACC CTATTAGTTG AAAGTTGAAA CTTTGTTCCC TACTCAATTC
CTAGTTGTGT AAATGTATGT ATATGTAATG CGTATAAAAC GTAGTACTTA
AATGACTAGG AGTGGTTCTT GAGACCGATG AGAGATGGGA GCAGAACTAA
AGATGATGAC ATAATTAAGA ACGAATTTGA AAGGCTCTTA GGTTTGAATC
CTATTCGAGA ATGTTTTTGT CAAAGATAGT GGCGATTTTG AACCAAAGAA
AACATTTAAA AAATCAGTAT CCGGTTACGT TCATGCAAAT AGAAAGTGGT
CTAGGATCTG ATTGTAATTT TAGACTTAAA GAGTCTCTTA AGATTCAATC
CTGGCTGTGT ACAAAACTAC AAATAATATA TTTTAGACTA TTTGGCCTTA
ACTAAACTTC CACTCATTAT TTACTGAGGT TAGAGAATAG ACTTGCGAAT
AAACACATTC CCGAGAAATA CTCATGATCC CATAATTAGT CAGAGGGTAT
GCCAATCAGA TCTAAGAACA CACATTCCCT CAAATTTAAA GACAGAAATT
TGTAGACTTT TTTTTGGCGT TAAAGGAAGA CTAAGTTTAT ACGTACATTT

TATTTTAAGT GGAAAACCGA AATTTTCCAT CGAAATATAT GAATTTAGTA
TATATATTTC TGCAATGTAC TATTTTGCTA TTTTGGCAAC TTTCAGTGGA
CTACTACTTT ATTACAATGT GTATGGATGC ATGAGTTTGA GTATACACAT
GTCTAAATGC ATGCTTTGCA AAACGTAACG GACCACAAAA GAGGATCCAT

5 GCAAATACAT CTCATAGCTT CCTCCATTAT TTTCCGACAC AAACAGAGCA.

10. The recombinant nucleic acid of claim 1 wherein the transcriptional regulatory region comprises (5' to 3'):

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CTGACTTC ACCAAAGAAA CAACTCGAGT CGTTATCCAT TCTCTACATC GCAACCCGGC CCAAACCGGT TTACCTCGTT GAGTACTCAT GCTACCTTCC ACCAACGCAT TGTAGATCAA GTATCTCCAA GGTCATGGAT 15 ATCTTTATC AAGTAAGAAA AGCTGATCCT TCTCGGAACG GCACGTGCGA TGACTCGTCG TGGCTTGACT TCTTGAGGAA GATTCAAGAA CGTTCAGGTC TAGGCGATGA AACTCACGGG CCCGAGGGGC TGCTTCAGGT CCCTCCCCGG AAGACTTTTG CGGCGGCGC TGAAGAGACG GAGCAAGTTA TCATTGGTGC GCTAGAAAAT CTATTCAAGA ACACCAACGT TAACCCTAAA GATATAGGTA 20 TACTTGTGGT GAACTCAAGC ATGTTTAATC CAACTCCATC GCTCTCCGCG ATGGTCGTTA ACACTTTCAA GCTCCGAAGC AACGTAAGAA GCTTTAACCT TGGTGGCATG GGTTGTAGTG CCGGCGTTAT AGCCATTGAT CTAGCAAAGG ACTTGTTGCA TGTCCATAAA AATACGTATG CTCTTGTGGT GAGCACAGAG AACATCACTT ATAACATTTA CGCTGGTGAT AATAGGTCCA TGATGGTTTC 25 AAATTGCTTG TTCCGTGTTG GTGGGGCCGC TATTTTGCTC TCCAACAAGC CTGGAGATCG TAGACGGTCC AAGTACGAGC TAGTTCACAC GGTTCGAACG CATACCGGAG CTGACGACAA GTCTTTTCGT TGCGTGCAAC AAGGAGACGA TGAGAACGGC AAAATCGGAG TGAGTTTGTC CAAGGACATA ACCGATGTTG CTGGTCGAAC GGTTAAGAAA AACATAGCAA CGTTGGGTCC GTTGATTCTT 30 CCGTTAAGCG AGAAACTTCT TTTTTTCGTT ACCTTCATGG GCAAGAAACT TTTCAAAGAT AAAATCAAAC ATTACTACGT CCCGGATTTC AAACTTGCTA TTGACCATTT TTGTATACAT GCCGGAGGCA GAGCCGTGAT TGATGTGCTA GAGAAGAACC TAGCCCTAGC ACCGATCGAT GTAGAGGCAT CAAGATCAAC GTTACATAGA TTTGGAAACA CTTCATCTAG CTCAATATGG TATGAGTTGG 35 CATACATAGA AGCAAAAGGA AGGATGAAGA AAGGTAATAA AGTTTGGCAG ATTGCTTTAG GGTCAGGCTT TAAGTGTAAC AGTGCAGTTT GGGTGGCTCT AAACAATGTC AAAGCTTCGA CAAATAGTCC TTGGGAACAC TGCATCGACA

GATACCCGGT CAAAATTGAT TCTGATTCAG GTAAGTCAGA GACTCGTGTC
CAAAACGGTC GGTCCTAATA AACGATGTTT GCTCTCTTTC GTTTCTTTT
ATTTGTTATA ATAATTTGAT GGCTACGATG TTTCTCTTGT TTGTTATGAA
TAAAGAATGC AATGGTGTTC TAGTATTTGA TTGTTTTACA TGTATGTATC
TCTTATTTAC ATGAAATTTT TAAACGCCTA AAAAAAAAA CGGAATTCCG.

11. The recombinant nucleic acid of claim 1 wherein the transcriptional regulatory region comprises (5' to 3'):

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5

CAGCTTAAC CGGTAAAATT GGCCTGTACA TATATTTACC ACTGAGTAAA GACATCAGTT AATGATTTGT TGTTACTCAA TTGGGCTAAG TGTATTATTA TATGTGTTGT ATATAATAAA GGTAGAACGT AAATTTACTA AGAATGTGTT TTTCCAATGT GATTGCTCTT 15 TGGCCTCTTA GGTTTGAATC CTACTCGAGA AGACTAATTT TAATTTACTG GCAAAAATAG AAATCAATTT ATAAGTGTTT AAACAAATCG ATGGTATAAC TGATTAGTGA TCACTCTTAG GTTTTGATCC AACTCGAGTA TTGAGTATTG AACGCTTTTT TTAAATAAAA TCTTGATTTT TAAATTGGTT TTTTGAGTAA AAAAGTTCTT AATATTTTCT CTTTGTTTTA ATGGGTTTGT TTTGCATTTT 20 ATAAGCTTAA TTTTTCTAAT TTAATATTTT ATCTATCATC GTCCGTAAAG TTTTATTTGG CACAAACTTG TTTTACTTTT CTACCTTATA ATTTGGGAAC TGGTTGAGTC AAAGCGTACC GGACAAATAT GTTTTATATT CTTATTTAAG AATTAACACT CATCTCATAA TTAGTCAGAG GCTAGGGAGA TTCAGCCAAT CAATGCTAAC AACAAAATTC TCTTAATGAT CTAACGATGC TATTTAATAT 25 TCGGATCAGT ATTCTTAAAT AAGAATATAA AACTAATTCA ATAGTTACAG ATAAAAACTT ATATAGACTT TTTTATTTGG AATATAAAAG TATCAATATA TTATAGACAA TATTTATAAC GTTAAAAATA CAATATTTAT ATTTTTTATA TATTTATTTC AAATTGAAAA GCATTACTTC TATCGAAATG AATTTTAGTA TATTAATTAA TATTTTTTTA ATCGGACTAC TTTCCTATTT TGGCACCTTT 30 CATCTGACTA CTAATTTATT TCAATGTGTA TGCATGCATG AGCATGAGTA ATACACATGT CTATATAAAT GCATGTAAAA CGTAACGGAC CACAAAAGTG GATCCATACA AATACATCTC ATCGCACCCT CTCCGACACA AAACTGAACA.

12. The recombinant nucleic acid of any one of claims 1 through 11 wherein the nucleic acid sequence encodes a translatable mRNA.

13. The recombinant nucleic acid of claim 12 wherein the nucleic acid sequence encodes an enzyme involved in lipid metabolism.

- 14. The recombinant nucleic acid of any one of claims 1 through 13, further comprising a transcription termination region operably linked to the nucleic acid sequence.
  - 15. A host cell comprising the recombinant nucleic acid of any one of claims 1 through 14.
  - 16. The host cell of claim 15, wherein the host cell is of a dicotyledonous plant species.

10

- 17. A plant comprising the recombinant nucleic acid of any one of claims 1 through 14.
- 18. The plant of claim 17, wherein the plant is of a dicotyledonous plant species.
- 15 19. A method of altering the phenotype of a seed comprising:
  - a) transforming a seed-bearing plant, or a progenitor of the seed-bearing plant, with a vector comprising the nucleic acid of any one of claims 1 through 14:
  - b) growing the seed-bearing plant to obtain seed under conditions wherein the nucleic acid sequence is expressed during embryogenesis under the control of the transcriptional regulatory region to alter the phenotype of the seed.

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20. A method of transforming a plant cell comprising transforming the plant cell with the recombinant nucleic acid of any one of claims 1 through 14.

25

Figure 1. Arabidopsis thaliana FAE1 promoter: (Length: 934 bp)

-950 ACTCA TAAAAACTAG TAGATTGGTT GGTTGGTTTC CATGTACCAG Atprofw > -900 AAGGCTTACC CTATTAGTTG AAAGTTGAAA CTTTGTTCCC TACTCAATTC -850 CTAGTTGTGT AAATGTATGT ATATGTAATG CGTATAAAAC GTAGTACTTA -800 AATGACTAGG AGTGGTTCTT GAGACCGATG AGAGATGGGA GCAGAACTAA -750 AGATGATGAC ATAATTAAGA ACGAATTTGA AAGGCTCTTA GGTTTGAATC -700 CTATTCGAGA ATGTTTTTGT CAAAGATAGT GGCGATTTTG AACCAAAGAA -650 AACATTTAAA AAATCAGTAT CCGGTTACGT TCATGCAAAT AGAAAGTGGT -600 CTAGGATCTG ATTGTAATTT TAGACTTAAA GAGTCTCTTA AGATTCAATC -550 CTGGCTGTGT ACAAAACTAC AAATAATATA TTTTAGACTA TTTGGCCTTA -500 ACTAAACTTC CACTCATTAT TTACTGAGGT TAGAGAATAG ACTTGCGAAT -450 AAACACATTC CCGAGAAATA CTCATGATCC CATAATTAGT CAGAGGGTAT -400 GCCAATCAGA TCTAAGAACA CACATTCCCT CAAATTTTAA TGCACATGTA -350 ATCATAGTTT AGCACAATTC AAAAATAATG TAGTATTAAA GACAGAAATT -300 TGTAGACTTT TTTTTGGCGT TAAAGGAAGA CTAAGTTTAT ACGTACATTT -250 TATTTTAAGT GGAAAACCGA AATTTTCCAT CGAAATATAT GAATTTAGTA -200 TATATATTC TGCAATGTAC TATTTTGCTA TTTTGGCAAC TTTCAGTGGA -150 CTACTACTTT ATTACAATGT GTATGGATGC ATGAGTTTGA GTATACACAT -100 GTCTAAATGC ATGCTTTGCA AAACGTAACG GACCACAAAA GAGGATCCAT -50 GCAAATACAT CTCATAGCTT CCTCCATTAT TTTCCGACAC AAACAGAGCA AtproRV 1 ATGACGTCCG TTAACGTTAA GCTCCTT

Figure 2. Brassica napus FAE1 promoter: (Length: 1588 bp)

-1600 GGTTGGGCAA ATCTGACTTC ACCAAAGAAA CAACTCGAGT CGTTATCCAT BnproFW → -1500 TCTCTACATC GCAACCGGC CCAAACCGGT TTACCTCGTT GAGTACTCAT -1450 GCTACCTTCC ACCAACGCAT TGTAGATCAA GTATCTCCAA GGTCATGGAT -1400 ATCTTTATC AAGTAAGAAA AGCTGATCCT TCTCGGAACG GCACGTGCGA -1350 TGACTCGTCG TGGCTTGACT TCTTGAGGAA GATTCAAGAA CGTTCAGGTC -1300 TAGGCGATGA AACTCACGGG CCCGAGGGGC TGCTTCAGGT CCCTCCCGG -1250 AAGACTTTTG CGGCGGCGCG TGAAGAGACG GAGCAAGTTA TCATTGGTGC -1200 GCTAGAAAAT CTATTCAAGA ACACCAACGT TAACCCTAAA GATATAGGTA -1150 TACTTGTGGT GAACTCAAGC ATGTTTAATC CAACTCCATC GCTCTCCGCG -1100 ATGGTCGTTA ACACTTTCAA GCTCCGAAGC AACGTAAGAA GCTTTAACCT -1050 TGGTGGCATG GGTTGTAGTG CCGGCGTTAT AGCCATTGAT CTAGCAAAGG -1000 ACTTGTTGCA TGTCCATAAA AATACGTATG CTCTTGTGGT GAGCACAGAG -950 AACATCACTT ATAACATTTA CGCTGGTGAT AATAGGTCCA TGATGGTTTC -900 AAATTGCTTG TTCCGTGTTG GTGGGGCCGC TATTTTGCTC TCCAACAAGC -850 CTGGAGATCG TAGACGGTCC AAGTACGAGC TAGTTCACAC GGTTCGAACG -800 CATACCGGAG CTGACGACAA GTCTTTTCGT TGCGTGCAAC AAGGAGACGA -750 TGAGAACGGC AAAATCGGAG TGAGTTTGTC CAAGGACATA ACCGATGTTG -700 CTGGTCGAAC GGTTAAGAAA AACATAGCAA CGTTGGGTCC GTTGATTCTT -650 CCGTTAAGCG AGAAACTTCT TTTTTTCGTT ACCTTCATGG GCAAGAAACT -600 TTTCAAAGAT AAAATCAAAC ATTACTACGT CCCGGATTTC AAACTTGCTA -550 TTGACCATTT TTGTATACAT GCCGGAGGCA GAGCCGTGAT TGATGTGCTA -500 GAGAAGAACC TAGCCCTAGC ACCGATCGAT GTAGAGGCAT CAAGATCAAC

### Figure 2 Continued Brassica napus FAE1 promoter:

-450	GTTACATAGA	TTTGGAAACA	CTTCATCTAG	CTCAATATGG	TATGAGTTG
-400	CATACATAGA	AGCAAAAGGA	AGGATGAAGA	AAGGTAATAA	AGTTTGGCAG
-350	ATTGCTTTAG	GGTCAGGCTT	TAAGTGTAAC	AGTGCAGTTT	GGGTGGCTCT
-300	AAACAATGTC	AAAGCTTCGA	CAAATAGTCC	TTGGGAACAC	TGCATCGACA
-250	GATACCCGGT	CAAAATTGAT	TCTGATTCAG	GTAAGTCAGA	GACTCGTGTC
-200	CAAAACGGTC	GGTCCTAATA	AACGATGTTT	GCTCTCTTTC	GTTTCTTTTT
-150	ATTTGTTATA	ATAATTTGAT	GGCTACGATG	TTTCTCTTGT	TTGTTATGA
-100	TAAAGAATGC	AATGGTGTTC	TAGTATTTGA	TTGTTTTACA	TGTATGTATC
-50	TCTTATTTAC	ATGAAATTTT			CGGAATTCC
			•	BnproRV	
1	ATGACGTCCA	TTAACGTAAA	GCTCCTTTAC	CATTACGTCA	TAACCAACCT
51	TTTCAACCTT		CGTTAACGGC	GATCGTCGCC	GGAAAAGCCI
	← Bnwalk				
101	ATCGGCTTAC	CATAGACGAT	CTTCACCACT	TATACTATTC	CTATCTCCAP
151	CACAACCTCA		TCCACTCTTT	GCCTTCACCG	
	<b>←</b>	Rowalki			

Figure 3. Lunaria annua FAE1 promoter: (Length: 1069 bp)

-1100 CG CCGGGGAGTT TCAGCTTAAC CGGTAAAATT LaproFW -1050 GGCCTGTACA TATATTTACC ACTGAGTAAA GACATCAGTT AATGATTTGT -1000 TGTTACTCAA TTGGGCTAAG TGTATTATTA TATGTGTTGT ATATAATAAA -950 GGTAGAACGT AAATTTACTA AGAATGTGTT TTTCCAATGT GATTGCTCTT -900 TGGCCTCTTA GGTTTGAATC CTACTCGAGA AGACTAATTT TAATTTACTG -850 GCAAAATAG AAATCAATTT ATAAGTGTTT AAACAAATCG ATGGTATAAC -800 TGATTAGTGA TCACTCTTAG GTTTTGATCC AACTCGAGTA TTGAGTATTG -750 AACGCTTTTT TTAAATAAAA TCTTGATTTT TAAATTGGTT TTTTGAGTAA -700 AAAAGTTCTT AATATTTTCT CTTTGTTTTA ATGGGTTTGT TTTGCATTTT -650 ATAAGCTTAA TTTTTCTAAT TTAATATTTT ATCTATCATC GTCCGTAAAG -600 TTTTATTTGG CACAAACTTG TTTTACTTTT CTACCTTATA ATTTGGGAAC -550 TGGTTGAGTC AAAGCGTACC GGACAAATAT GTTTTATATT CTTATTTAAG -500 AATTAACACT CATCTCATAA TTAGTCAGAG GCTAGGGAGA TTCAGCCAAT -450 CAATGCTAAC AACAAAATTC TCTTAATGAT CTAACGATGC TATTTAATAT -400 TCGGATCAGT ATTCTTAAAT AAGAATATAA AACTAATTCA ATAGTTACAG -350 ATAAAAACTT ATATAGACTT TTTTATTTGG AATATAAAAG TATCAATATA -300 TTATAGACAA TATTTATAAC GTTAAAAATA CAATATTTAT ATTTTTTATA -250 TATTTATTC AAATTGAAAA GCATTACTTC TATCGAAATG AATTTTAGTA -200 TATTAATTAA TATTTTTTTA ATCGGACTAC TTTCCTATTT TGGCACCTTT -150 CATCTGACTA CTAATTTATT TCAATGTGTA TGCATGCATG AGCATGAGTA

## Figure 3 Continued. Lunaria annua FAE1 promoter:

- -100 ATACACATGT CTATATAAAT GCATGTAAAA CGTAACGGAC CACAAAAGTG
- -50 GATCCATACA AATACATCTC ATCGCACCCT CTCCGACACA AAACTGAACA

  ← Laprorv
  - 1 ATGACGTCTG TGAACGTAAA ACTCCTTTAC CATTACGTCA TAACCAACTT
- 51 TTTCAACCTC TGTTTCTTCC CACTGACGGG GATCCTCGCC GGAAAAGGCT
- 101 CTCGTCTTAC CACAAACGAT CTCCACCA

Figure 4: Alignment of A.t., L.a. and B.n. FAE1 promoters

CLUSTAL W (1.74) multiple sequence alignment

A.t.	
L.a.	
В.п.	GGTTGGGCAAATCTGACTTCACCAAAGAAACAACTCGAGTCGTTATCCATCTCCTCATAA
A.t.	
L.a.	
B.n.	CCATCGCTCCACTCTTTGCCTTCACCGTTTTCGGTTCGG
A.t.	
L.a.	
B.n.	CCAAACCGGTTTACCTCGTTGAGTACTCATGCTACCTTCCACCAACGCATTGTAGATCAA
A.t.	
L.a.	
B.n.	GTATCTCCAAGGTCATGGATATCTTTTATCAAGTAAGAAAAGCTGATCCTTCTCGGAACG
A. t.	
L.a.	
B.n.	GCACGTGCGATGACTCGTCGTGGCTTGACTTCTTGAGGAAGATTCAAGAACGTTCAGGTC
A.t.	
L.a.	
B.n.	TAGGCGATGAAACTCACGGGCCCGAGGGGCTGCTTCAGGTCCCTCCC
A.t.	
L.a.	
B.n.	CGGCGGCGCGTGAAGAGACGGAGCAAGTTATCATTGGTGCGCTAGAAAATCTATTCAAGA
A.t. L.a.	
B.n.	ACACCAACGTTAACCCTAAAGATATAGGTATACTTGTGGTGAACTCAAGCATGTTTAATC
A.t.	
L.a.	CGCCGGGGAGTTTCAGCTTAA
B.n.	CAACTCCATCGCTCTCCGCGATGGTCGTTAACACTTTCAAGCTCCGAAGCAACGTAAGAA
Con. 4	инииниининининининининининининининининин
A.t.	actcataaaactagtagattggttgg
L.a.	CCGGTAAAATTGGCCTGTACATATTTTACCACTGAGTAAAGACATCAGTTAATGATTTG
B.n.	GCTTTAACCTTGGTGGCATGGGTTGTAGTGCCGGCGTTATAGCCATTGATCTAGCAAAGG
Con. 4	NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNSKSRKWTWARMYCKYRRWYNNKSRWWKG

## Figure 4 Continued: Alignment of A.t., L.a. and B.n. FAE1 promoters

A.t. L.a. B.n.	TTGGTTTCCATGTACCAGAAGGCTTACCCTAT-TAGTTGAAAGTTGAAACTTTGTTCC TTGTTACTCAATTGGGCTAAGTGTATTATTATAT-GTGTTGTATATAATAAAGGTAGAAC ACTTGTTGCATGTCCATAAAAATACGTATGCTCTTGTGGTGAGCACAGAGAACATCACTT
Con.4	WYKKKWYBCANNTSBRYHARRWKDMKTAYBMTMTNKWGKTGWRHRYWRWRAMBDTVDHHY
A.t. L.a.	CTACTCAATTCCTAGTTGTGTAAATGTATGTATATGTAATGCGTATAAAACGTA GTAAATTTACTAAGAATGTGTTTTTCCAATGTGATTGCTCTTTTGGCCTCTTAGGTTTG ATAA-CATTTACGCTGGTGATAATAGGTCCATGATGGTTTCAAATTGCTTGTTCCGTGTT
B.n.	** ** ** * * * * * * * * * * * * * * *
Con.4	VTAMNNAWTTMCMMDKDDKRTRWWWKKNNNATGWDDDTKYHMWNNNGCBTVTWMVRYKTD
A.t.	GTACTTAAATGACTAGGAGTGGTTCTTGAGACCGATGAGAGATGGGAG-CAGAACTAAAG
L.a.	AATCCTACTCGAGAAG-ACTAATTTTAATTTACTGGCAAAAATAGAAA-TCAATTTATAA
B.n.	GGTGGGGC-CGCTATTTTGCTCTCCAACAAGCCTGGAGATCGTAGACGGTCCAAGTACGA
Con.4	RDWSBKRMNYGMBWWKNWSYDVTYYWWVWDDMCKRKVRRWVRTRGRMRNYMVAWBTAHRR
A.t.	ATGATGACATAATTAAGAACGAATTTGA-AAGG-CTCTTAGGTTTGAATCCT
L.a.	GTGTTTAAACAAATCGATGGTATAACTGATTAGT-GATCACTCTTAGGTTTTGATCCA
B.n.	GCTAGTTCACACGGTTCGAACGCATACCGGAGCTGACGACAAGTCTTTTCGTTGCGTGCA
Con.4	RYNNGWTBAMAYRRWTMNNNNNAKAMCKRAKYWGWNRABVNSTCTTWKSKTTKVRTSCW
A.t.	ATTCGAGAATGTTTTTGTCAAAGATAGTGGCGATTTTGAACCAAAGAAAACATTTAAA-A
L.a.	ACTCGAGTATTGAGTATTGAACGCTTTTTTTAAATAAAATCTTGATTTTTA-A
B.n.	ACAAGGA-GACGATGAGAACGGCAAAATCGGAGTGAGTTTGTCCAAGGACATA * ** ** ** ** ** ** ** ** ** ** ** ** *
Con.4	ANNCRAGDANKDHKWWKWSAAMGVYWNNNNNNWTYKKARHBARWDWVWHSAWKKWHANA
A.t.	AATCAGTATCCGGTTACGTTCATGCAAATAGAAAGTGGTCTAGGATCTGATT-
L.a.	ATTGGTTTTTTGAGTAAAAAGTTCTTAATATTTTCTCTTTGTTTTAATGGGTTTGTTT
B.n.	ACCGATGTTGCTGGTCGAACGGTTAAGAAAACATAGCAACGTTGGGTCCGTTGA
Con.4	AHYSRKKWTBYKRKTMVNNNNGTTMWKRMWAWYWKMDMDWBGTYNNNNNGGRTYYGWTKN
A.t.	GTAATTTTAGACTTAAAGAGTCTCTTAAGATTCAATCCTGGCT-GTGTACAAAACT
L.a.	TGCATTTTATAAGCTTAATTTTCTAATTTAATATTTTATCTATC
B.n.	TTCTTCCGTTAAGCGAGAAACTTCTTTTTTTCGTTACCTTCATGGGCAAGAAACTT
Con.4	KKMWTYYKWKANNCKWRAWDHKTCTHNNTTWWKMKTYWNNCYWKSMTNGKSHRBAAAVYT
A.t.	ACAAATAATATATTTTTAGACTATTTGGCCTTAACTAAACTTCCA-CTCATTATTTA
L.a.	TTATTTGGCACAAACTTGTTTTACTTTTCTACCTTATAATTTGGGAACTGGTTGAGT-
B.n.	TTCAAAGATAAAATCAAACATTACTACGTCCCGGATTTCAAACTTGCTATTGACCATTTT
Con.4	WYMWWRRYAHANNNWDYWWKACTWYKYBVCSKWWNNYAAWYTKSSWNYTSRYYRWKTN
A.t.	-CTGAGGTTAGAGAATAGACTTGCGAATAAACACATTCCCGAGAAATACTCATGATCC
L.a.	-CAAAGCGTACCGGACAAATATGTTT-TATATTCTTATTTAAGAATTAACACTCATCT
B.n.	TGTATACATGCCGGAGGCAGAGCCGTGATTGATGTGCTAGAGAAGAACCTAGCCCTAGCA
Con.4	NSWRWRSDTRSMGRANNYARABHYGYKWNTRWWBWSHTWBHBRAGAAHYWMBMMYBAKCH

Figure 4 Continued: Alignment of A.L., L.a. and B.n. FAE1 promoters

	•
2 4	CE3
A.t.	CATAATTAGTCAGAGGGTATGCCAATCAGATCTAAGAACACATTCCCTC
L.a.	CATAATTAGTCAGAGGCTAGGGAGATTCAGCCAATCAATGCTAACAACAAA-ATTCTCTT
B.n.	CCGATCGATGTAGAGGCATCAAGATCAACGTTACATAGATTTGG
Con.4	CMKAWYKAKKYAGAGGSNNNNNNNNNNNNNNNNNATCARDDYYAASRWYAMANAKWYYYKB
A.t.	AAATTTTAATGCAC-ATGTAATCATAGTTTAGCACAATTCAAAA
L.a.	AATGATCTAACGATGCTATTTAATATTCGGATCAGTATTCTTAAATAAGAATATAAAA
B.n.	AAACACTTCATCTAGCTCAATATGGTATGAGTTGGCATACAT-AGAAG-CAAAA
Con.4	AANNAYYTHANNWWGCWNNATDTRRTMWKNNNNNNAGTWKNNNNNNAKNASAAKNYAAAA
A.t.	ATAATGTAGTA-TTAAAGACAGAAATTTGTAGACTTTTTTTTGGCGT-TAAAGG
L.a.	CTAATTCAATAGTTACAGATAAAAACTTATATAGACTTTTTTATTTGGAATATAAAAG
B.n.	GGAAGGATGAA-GAAAGGTAATAAAGTTTGGCAGATTGCTTTAGGGTCAGGCTTTAAGTG
Con.4	VKAAKKHWRWANKWAMRGWHADAAABTTDKRNNGAYTKYTTTNNNNTYRGVVTNTAARDG
A.t.	AAGACTAAGTTTATA-CGTACATTT-TATTTTAAGT
L.a.	TATCAATATATATA-GACAATATTTATAACGTTAAAAATACAATATTTATATTTTTAT
B.n.	TAACAGTGCAGTTTGGGTGGCTCTAAACAATGTCAAAGCTTCGACAAATAGTCCTTGGGA
Con.4	WANNNNNNNNNNNNGWSDMWVTWWAYANYGTNNNNNNNNNNAYAWWTNKWYYTTDDRW
	CE1
A.t.	GGAAAACCGAAATTTTCCATCGAAATATATGAATTT-AGTATAT
L.a.	ATATTTATTTCAAATTGAAAAGCATTACTTCTATCGAAATGAATTTTAGTATATTAAT
B.n.	ACACTGCATCGACAGATACCCGGTCAAAATTGATTCTGATTCAGGTAAGTCAGA
Con.4	RBAYTNNNNNRMAYYGAYADDYAYYMSDTCDAWMKWDATKMNNATTYNRGTAWRTNNNN
7 6	G-box2
A.t.	ATATTTCTGCAATGTACTATTTTGCTATTTTGGCAA-CTTTCAGTGGACTAC
L.a.	TAATATTTTTTAATCGGACTACTTTCCTATTTTGGCAC-CTTTCATCTGACTAC
B.n.	GACTCGTGTCCAAAACGGTCGGTCCTAATAAACGATGTTTGCTCTCTTTTCGTTTCTTT
Con.4	NNMTMKTKYYBHAAWNNNNNGKMCTAHTWWVCKATKTTKGCWMNCTTTCRKYKNNCTWY
3 4	G-box1
A.t.	TACTTTATTACAATGTGTATGGATGC-ATGAGTTTGAGTA-TACACATGTCTAAA
L.a.	TAATTTATTTCAATGTGTATGCATGC-ATGAGCATGAGTAATACACATGTCTATA
B.n.	TTATTTGTTATAATAATTTGATGGCTACGATGTTTCTCTTTGTTTG
Con.4	TWMTTTRTTWYAATRWKTNNATGSMTRCNATGWKNNNYWTGWKTRWTAYRMATRWMKAWW
_	A-300 EM1 ABA
A.t.	TGCATGCT-TTGCAAAACGTAACGGACC-ACAAAAGAGGGATCCA <u>TGCAAAT</u> ACATCTCAT
L.a.	TAAATGCA-T-GTAAAACGTAACGGACC-ACAAAAGTGGATCCATACAAATACATCTCAT
B.n.	GCAATGGTGTTCTAGTATTTGATTGTTTTACATGTATCTCTT-ATTTACATGAAAT
Con.4.	KVMATGSWNTNSYARWAYKTRAYKGWYYNACAWRWRWGKATCYMTDNAWWTACATSWMAT
A.t.	AGC-TTCCTCCATTATTTTCCGACACAAA-CAGAGCA
L.a.	CGC-ACCCTCTCCGACACAAAACTGAACA
B.n.	TTTTAAACGCCTAAAAAAAAAACGGAATTCCG
Con.4	HKYNWHMCKCNNNNNNNTMMRAMAMAANCDGARYWNNN

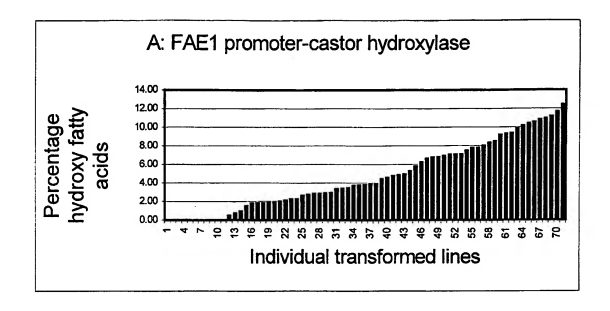
# Figure 5: Alignment of A.t. and L.a. FAE1 promoters CLUSTAL W (1.74) multiple sequence alignment

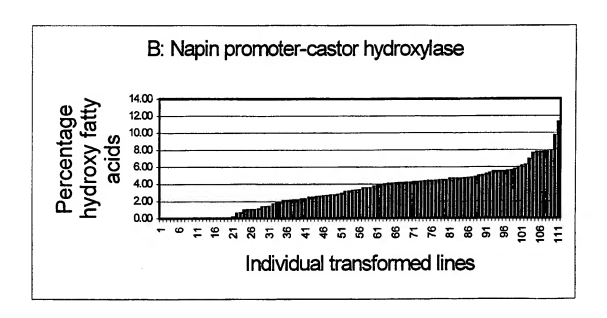
A.t. L.a.	CGCCGGGGAGTTCAGCTTAACCGGTAAAATTGGCCTGTACATATATTTACCACTGAGTA
	*** * *
Con.5	ACTSAKWA
A.t.	AAACTAGTAGATTGGTTGGTTGGTTTCCATGTACCAGAAGGCTTACCCTATTAGTT
L.a.	AAGACATCAGTTAATGATTTGTTGTTACTCAATTGGGCTAAGTGTATTATTATATGTGTT ** * * * * * * * * * * * * * * * * *
Con.5	AARMYAKYAGWTNNTGRTTKGTTGKTWYYCANNTGKRCYARRWGKMTTAYYMTATKWGTT
A.t.	GAAAGTTGAAACTTTGTTCCCTACTCAATTCCTAGTTGTGTAAATGTATGTATATGT
L.a.	GTATATAATAAAGGTAGAACGTAAATTTACTAAGAATGTGTTTTTCCAATGTGATTGC
Con.5	GWAWRTWRWAAMKKTRKWMCSTAMNNAWTTMCTARKWRTGTRWWTKTNNNATGTRWWTGY
A.t.	AATGCGTATAAAACGTAGTACTTAAATGACTAGGAGTGGTTCTTGAGACCGATGAGA
L.a.	TCTTTGGCCTCTTAGGTTTGAATCCTACTCGAGAAG-ACTAATTTTAATTTACTGGCAAA
Con.5	WMTNNNGCSTMTWARRYKTRRWWCYTAMWYGASWAGNASTRRTTYTWRWKWMCKRKSARA
A.t.	GATGGGAGCAGAACTAAAGATGATGACATAATTAAGAACGAATTTGAAAGG-CT
L.a.	AATAGAAATCAATTTATAAGTGTTTAAACAAATCGATGGTATAACTGATTAGTGATCACT
Con.5	RATRGRARYMRAWYTAWARRTGWTKAMAYAAWTMNNNNNAKAACKRATTWGWRAKSNCT
A. t.	CTTAGGTTTGAATCCTATTCGAGAATGTTTTTGTCAAAGATAGTGGCGATTTTGAACCAA
L.a.	CTTAGGTTTTGATCCAACTCGAGTATTGAGTATTGAACGCTTTTTTTTAAATAA
Con.5	CTTAGGTTTKRATCCWAYTCGAGWATKKWKTWKTSAAMGMTWNNNNNNTTTTKAAMYAA
A.t.	AGAAAACATTTAAAAAATCAGTATCCGGTTACGTTCATGCAAATAGAAAGTGGTCT
L.a.	AATCTTGATTTTTAAATTGGTTTTTTGAGTAAAAAAGTTCTTAATATTTTCTCTTTGTTT
Con.5	ARWMWWSATTTWWAAAWTSRKTWTYYGRKTAMNNNNGTTCWTRMWAWTWKMWMKTKGTTT
A.t.	AGGATCTGATTGTAATTTTAGACTTAAAGAGTCTCTTAAGATTCAATCCTGGC
L.a.	TAATGGGTTTGTTTTGCATTTTATAAGCTTAATTTTTCTAATTTAATATTTTATCTATC
Con.5	WNNNGGRTYTGWTTKKMATTTTAKANNCTTAAWKWKTCTMNNTTAAKATTYWATCYWKSM
A.t.	T-GTGTACAAAACTACAAATAATATATTTTAGACTATTTGGCCTTAACTAAACTTC
L.a.	TCGTCCGTAAAGTTTTATTTGGCACAAACTTGTTTTACTTTTCTACCTTATAATTTGG
Con.5	TNGTSYRYAAARYTWYAWWTRRYAYANNNNTKTTWKACTWTTYKRCCTTANNTAAWYTKS
A.t.	CA-CTCATTATTTACTGAGGTTAGAGAATAGACTTGCGAATAAACACATTCCCGAGAAAT
L.a.	GAACTGGTTGAGT-CAAAGCGTACCGGACAAATATGTTT-TATATTCTTATTTAAGAATT
Con.5	SANCTSRTTRWKTNCWRAGSKTASMGRAYARAYWTGYKWNTAWAYWCWTWYYYRAGAAWT
	-400
A.t432 L.a.	ACTCATGATCCCATAATTAGTCAGAGGGTATG
Con 5	*
Con.5	AMYMMTSATCYCATAATTAGTCAGAGGSTAKGNNNNNNNNCCAATCARWKCTAASAACA

## Figure 5 Continued: Alignment of A.t. and L.a. FAE1 promoters

A.t. L.a.	-381	CACATTCCCTCAAATTTTAATGCACATGTAATCATAGTTTAGCA AA-ATTCTCTTAATGATCTAACGATGCT-ATTTAATATTCGGATCAGTATTCTTAAATAA
Con.5		MANATTCYCTYAANNATYTWANNATGCWNATKTAATMWTNNNNNAGTWTNNNNNNAKMA
A.t. L.a.	-337	CAATTCAAAAATAATGTAGTA-TTAAAGACAGAAATTTGTAGACTTTTTTTTGGCG GAATATAAAACTAATTCAATAGTTACAGATAAAAACTTATATAGACTTTTTTATTTGGAA
Con.5		SAATWYAAAAMTAATKYARTANTTAMAGAYARAAAYTTRTANNGACTTTTTTNNTTGGMR
A.t. L.a.	-282	T-TAAAGGAAGACTAAGTTTATA-CGTACATTT-TAT TATAAAAGTATCAATATTATATAGACAATATTTATAACGTTAAAAATACAATATTTATAT
Con.5		TNTAAARGWANNNNNNNNNNNNNGACWAWRTTTATANCGTNNNNNNNNNNNNAYATTTNTAT
A.t. L.a.	-247	TTTAAGTGGAAAACCGAAATTTTCCATCGAAATATATGAATTT-AGTATA TTTTTATATATTTTATTT
Con.5		TTTWWRTRKANNNNNNNAAAYYGAAAWKNNTTMCWTCKAWMKAWATGAATTTNAGTATA
A.t. L.a.	-198	TATATTTCTGCAAT-GTACTATTTTGCTATTTTTGCAACTTTCAGTGGACTACT TTAATTAATATTTTTTTTAATCGGACTACTTTCCTATTTTGGCACCTTTCATCTGACTACT
Con.5		TNNNNNATATTTYTKYAATNGKACTAYTTTSCTATTTTGGCAMCTTTCAKYKGACTACT
A.t. L.a.	-145	ACTTTATTACAATGTGTATGGATGCATGAGTTTGAGTA-TACACATGTCTAAATGCATGC AATTTATTTCAATGTGTATGCATGCATGAGCATGAGTAATACACATGTCTATATAAATGC
Con.5		AMTTTATTWCAATGTGTATGSATGCATGAGYWTGAGTANTACACATGTCTAWATRMATGC
A.t. L.a.	-86	TTTGCAAAACGTAACGGACCACAAAAGAGGATCCATGCAAATACATCTCATAGCTTCCTCAT-GTAAAACGTAACGGACCACAAAAGTGGATCCATACAAATACATCTCATCGCACCCTC
Con.5		WTNGYAAAACGTAACGGACCACAAAAGWGGATCCATRCAAATACATCTCATMGCWYCCTC
A.t. L.a.	-26	CATTATTTTCCGACACAAA-CAGAGCATCCGACACAAAACTGAACA
Con.5		NNNNNNTCCGACACAAANCWGARCA

Figure 6





## Figure 7: Alignment of B.n. and L.a. FAEI promoters

CLUSTAL W (1.81) multiple sequence alignment

BnFAEl LaFAEl	GGTTGGGCAAATCTGACTTCACCAAAGAAACAACTCGAGTCGTTATCCATCTCCTCATAA	60
BnFAEl LaFAEl	CCATCGCTCCACTCTTTGCCTTCACCGTTTTCGGTTCTGCTACATCGCAACCCGGC	120
BnFAE1 LaFAE1	CCAAACCGGTTTACCTCGTTGAGTACTCATGCTACCTTCCACCAACGCATTGTAGATCAACGCCGGGGAGT-TTCAGCTTAACCGGTAAAATTGGCCTGTACATATA	180 46
BnFAE1 LaFAE1	GTATCTCCAAGGTCATGGATATCTTTTATCAAGTAAGAAAAGCTGATCCTTCTCGGAACG TTTACCACTGAGT-AAAGACATCAGTTAATGATTTGTTGTTACTCAATTGGGCT	240 99
BnFAE1 LaFAE1	GCACGTGCGATGACTCGTCGTGGCTTGACTTCTTGAGGAAGATTCAAGAACGTTCAGGTCAGGTCAAGTGTATTATTATATGTGTTGTATATAATAAAGGTAGAACGT-AAATT	300 147
BnFAE1 LaFAE1	TAGGCGATGAAACTCACGGGCCCGAGGGGCTGCTTCAGGTCCCTCCC	360 205
BnFAE1 LaFAE1	CGGCGGCGCGTGAAGAGACGGAGCAAGTTATCATTGGTGCGCTAGAAAATCTATTCAAGA CTCGAGAAGACTAATTTTAAT-TTACTGGCAAAAATAGAAATCAATTTATAA	420 256
BnFAE1 LaFAE1	ACACCAACGTTAACCCTAAAGATATAGGTATACTTGTGGTGAACTCAAGCATGTTTAATC GTGTTTAAACAAATCGATGGTATAACTG-ATTAGTGATCACTCTTAGGTTTTGATC	480 311
BnFAEl LaFAEl	CAACTCCATCGCTCTCCGCGATGGTCGTTAACACTTTCAAGCTCCGAAGCAACGTAAGAA SCAACTCGAGTATTGAACGCTTTTTTTAAATAAAATCTTGA	540 358
BnfAE1 LafAE1	GCTTTAACCTTGGTGGCATGGGTTGTAGTGCCGGCGTTATAGCCATTGATCTAGCAAAGG 6 TTTTTAAA-TTGGTTTTTTGAGTAAAAAAGTTCTTAATATTTTCTCTT-TGTTTTAATGG 4	500 116
BnFAE1 LaFAE1	ACTTGTTGCATGTCC-ATAAAAATACGTATGCTCTTGTGGTGAGCACAGAGAACATCACT 6 GTTTGTTTTGCATTTTATAAGCTTAATTTTCTAATTTAAT-ATTTTATCTATCATCGTC 4	59 75
BnFAEl LaFAEl	TATAACATTTACGCTGGTGATAATAGGTCCATGATGGTTTCAAATTGCTTGTTCCGTGTT 7 CGTAAAGTTTTATTTGGCACAAACTTGTTTTACTTTTCTACCTTATA 5	19 22
BnFAEl LaFAEl	GGTGGGGCCGCTATTTTGCTCTCCAACAAGCCTGGAGATCGTAGACGGTCCAAGTACGAG 7 ATTTGGGA-ACTGGTTGAGTCAAAGCGTACCGGACAAATATGTTTTATATTC 5	79 73
BnFAE1 LaFAE1	CTAGTTCACACGGTTCGAACGCATACCGGAGCTGACGACAAGTCTTTTCGTTGCGTGCAA 8:	39 24
BnFAE1 LaFAE1	CAAGGAGACGATGAGAACGGCAAAATCGGAGTGAGTTTGTCCAAGGACATAACCGATGTT 89 CAGCCAATCAATGCTAACAACAAAATTCTCTTAATGATCTAACGATGCTATTTAATAT 68	99 82

## Figure 7 Continued: Alignment of B.n. and La. FAE1 promoters

LaFAEl	TCGGATCAGTATTCTTAAATAAGAATATAAAACTAATTCAATAGTTACAG	
	* * * * * * * * * * * * * * * * * * * *	
BnFAE1	CGAGAAACTTCTTTTTTCGTTACCTTCATGGGCAAGAAACTTTTCAAAGATAAAATCAA	1018
LafAEl	ATAAAAACTTATATAGACTTTTTTATTTG-GAATATAAAAGTATCAATATATTATAGACA	
BnFAEl	ACATTACTACGTCCCGGATTTCAAACTTGCTATTGACCATTTTTGTATACATGCCGGAGG	1078
LaFAEl	ATATTTATAACGTTAAAAATACAATATTTATATTTTTTATATTTTTTCAAA	345
BnFAE1	CAGAGCCGTGATTGATGTGCTAGAGAAGAACCTAGCCCTAGCACCGATCGAT	1138
LaFAE1	TTGAAAAGCATTACTTCTATCGAAATGAATTTTAGTATATTAATTAATTTTTTTT 9	
BnFAE1	ATCAAGATCAACGTTACATAGATTTGGAAACACTTCATCTAGCTCAATATGGTATGAGTT 1	1198
LaFAEl	AATCGGACTACTTTCCTATTTTGGCACCTTTCATCTGACTACT	
BnFAEl	GGCATACATAGAAGCAAAAGGAAGGATGAAGAAGGTAATAAAGTTTGGCAGATTGCTTT 1	258
LaFAE1	AATTTATTCAATGTGTATGCATGCATGAGCATGAGTAATACACATGTCTAT	
BnFAE1	AGGGTCAGGCTTTAAGTGTAACAGTGCAGTTTGGGTGGCTCTAAACAATGTCAAAGCTTC 1	.318
Lafae1	ATAAATGCATGTAAAACGTAACGG-ACCACAAAAGTGGATCCATACAAATACATCTCATC 1	.055
Bn <b>FA</b> El	GACAAATAGTCCTTGGGAACACTGCATCGACAGATACCCGGTCAAAATTGATTCTGATTC 1	.378
LafAE1	G-CACCCTCTCCGACACAAAACTGAACA	082
BnFAE1	AGGTAAGTCAGAGACTCGTGTCCAAAACGGTCGGTCCTAATAAACGATGTTTGCTCTCTT 1	438
Lafael		
BnFAE1	TCGTTTCTTTTATTTGTTATAATAATTTGATGGCTACGATGTTTCTCTTGTTTGT	100
LafAE1		430
BnFAE1	AATAAAGAATGCAATGGTGTTCTAGTATTTGATTGTTTTACATGTATGT	558
LafAEl		•
BnFAE1	ACATGAAATTTTTAAACGCCTAAAAAAAAAAAAACGGAATTCCG 1600	
aFAE1	2000	

## Figure 8: Alignment of B.n. and A.t. FAE1 promoters

CLUSTAL W (1.81) multiple sequence alignment

	•	
AtFAE1		
BnFAE1	GGTTGGGCAAATCTGACTTCACCAAAGAAACAACTCGAGTCGTTATCCATCTCCTCATAA	60
		•
AtFAE1		
BnFAE1	CCATCGCTCCACTCTTTGCCTTCACCGTTTTCGGTTCGG	120
AtFAE1	= = = = = = = = = = = = = = = = = = =	
Bn <b>FAE</b> l	CCAAACCGGTTTACCTCGTTGAGTACTCATGCTACCTTCCACCAACGCATTGTAGATCAA	180
AtFAE1	ACT CATARAA	
BnFAEl	GTATCTCCAAGGTCATGGATATCTTTTATCAAGTAAGAAAAGCTGATCCTTCTCGGAACG	240
	*** **	
145151		
AtFAEL	ACTAGTAGATTGGTTGGTTGGTTTCCATGTACCAGAAGGCTTACCCTATTAGT	63
BnFAE1	GCACGTGCGATGACTCGTCGTCGCTTGACTTCTTGAGGAAGATTCAAGAACGTTCAGGTC	300
ALFAE1	TChilicatch incar accompany of the contract of	
BnFAEl	TGAAAGTTGAAACTT-TGTTCCCTACTCAATTCCTAGTTGTGTAAATGTATGTATATG	120
2111111111	TAGGCGATGAAACTCACGGGCCCGAGGGGCTGCTTCAGGTCCCTCCC	160
AtFAE1	TAATG-CGTATAAAACGTAGTACTTAAATGACTAGGAGTGGTTCTTGAGACCGATGAGAG	70
BnFAE1	CGGCGGCGCGTGAAGAGAGGGC-AAGTTATCATTGGTGCGCTAGAAAATCTATTCAAG	110
	* ** * ** * * * * * * * * * * * * * *	113
AtFAE1	ATGGGAGCAGAACTAAAGATGATGACATAATTAAGAACGAATTTGAAAGGCTCTTA 2	75
BnFAE1	AACACCAACGTTAACCCTAAAGATATAGGTATACTTGTGG-TGAACTCAAGCATGTTTAA 4	78
	* * ******* * *** * * * * * * * * * *	
AtFAE1	GGTTTGAATCCTATTCGAGAATGTTTTTGTCAAAGATAGTGGCGA-TTTTGAACCAAAGA 2	94
BnFAE1	TCCAACTCCATCGCTCTCCGCGATGGTCGTTAACACTTTCAAGCTCCGAAGCAACGT 5	35
	* ** * ** * * * * * * * * * * * * * * *	
AtFAE1	AAACATTTAAAAAATCAGTATCCGGTTAC-GTTCATGCAA-ATAGAAAGTGGTCTAGG 3	50
BnFAEl	AAGAAGCTTTAACCTTGGTGGCATGGGTTGTAGTGCCGGCGTTATAGCCATTGATCTAGC 5	95
	** * * * * * * * * * * * * * * * * * * *	
AtFAE1	ATCTGATTGTAATTTTAGACTTAAAGAGTCTCTTAAGATTCAATCCTGGCTGTGTACAAA 4	10
BnFAE1	AAAGGACTTGTTGCATGTCCATAAAAATACGTATGCTCTTGTGGTGAGCACAGAGAAC 6	53
	* ** ** * * * * * * * * * * * * * * * *	
AtFAE1		
BnFAE1	ACTACAAATATAT——TTTAGACTATTTGGCCTTAACTAAACTTCCACTCATTATTT 4	67
DIILINGI	ATCACTTATAACATTTACGCTGGTGATAATAGGTCCATGATGGTTTCAAATTGCTTGTTC 7	13
AtFAE1	ACTGAGGTTAGAGA-ATAGACTTGCGAATAAACACATTCCCGAGAAATACTCATGATCCC 5	26
BnFAE1	CGTGTTGGTGGGGCCGCTATTTTGCTCTCCAACAAGCCTGGAGATCGTAGACGGTCCA 7	
	** * * * * * * * * * * * * * * * * * * *	, ±
	CE3	
AtFAE1	ATAATTAGTCAGAGGGTATGCCAATCAGATCTAAGAACACATTCCCTCAAATTTTA 5	84
BnFAEl	AGTACGAGCTAGTTCACACGGTTCGAACGCATACCGGAGCTGACGACAAGTCTTTTCGTT 83	
	* * ** ** * * * * * * * * * * * * *	_
At <b>FAE</b> l	ATGCACATGTAATCATAGTTTAGCACAATTCAAAAATAATGTAGTATTAAAGACAGAAAT 64	
BnFAE1	GCGTGCAACAAGGAGACGATGAGAACGGCAAAATCGGAGTGAGT	
	* ** * * * * * * * * * * * * * * * * * *	

## Figure 8 Continued: Alignment of B.n. and A.t. FAE1 promoters

ACLAET	TTGTAGACTTTTTTTTGGCGTTAAAGGAAGACTAAGTTTATACGTACATTTTAT 698
BnFAE1	CCGATGTTGCTGGTCGAACGGTTAAGAAAAACATAGCAACGTTGGGTCCGTTGATTCTTC 951
	* * * * * * * * * * * * * * * * * * * *
AtFAE1	T-TTAAGTGGAAAACCGAAATTTTCCATCGAAATATATGAATTTAGTATATATA 751
BnFAE1	CGTTAAGCGAGAAACTTCTTTTTTCGTTACCTTCATGGGCAAGAAACTTTTCAAAGATA 1013
	***** * *** *** * * * * * * * * *
	G box 2
AtFAE1	TTTCTGCAATGTACTATTTTGCTATTTTGGCAACTTTCAGTGGACTACTACTTTAT-TAC 810
BnFAE1	AAATCAAACATTACTACGTCCCGGATTTCA-AACTTGCTATTGACCATTTTTGTATACAT 1070
	* **** * * *** **** * * *** * * *** *
AtFAE1	G-box 1
BnFAE1	AATGTGTATGGATGCATGAGTT-TGAGTATACACATGTCTAAATGCATGCTTTGCAAAAC 869
BULAET	GCCGGAGGCAGAGCCGTGATTGATGTGCTAGAGAAGAACCTAGCCCTAGCACCGATCGAT
	* ** * *** * * * * * * * * *
AtFAE1	GTAACGG-ACCACAAAAGAGGATCCATGCAAATACATCTCATAGCTTCCTCCAT 922
BnFAE1	GTAGAGGCATCAAGATCAACGTTACATAGATTTGGAAACACTTCATCTAGCTCAATATGG 1190
	*** ** * * * * * * * * * * * * * * * * *
AtFAEl	TATTTTCCGACACAAACAGA-GCA945
BnFAE1	TATGAGTTGGCATACATAGAAGCAAAAGGAAGGATGAAGAAAGGTAATAAAGTTTGGCAG 1250
	***
AtFAE1	
BnFAE1	ATTGCTTTAGGGTCAGGCTTTAAGTGTAACAGTGCAGTTTGGGTGGCTCTAAACAATGTC 1310
	***************************************
145351	
AtFAEl BnFAEl	
DREAL	AAAGCTTCGACAAATAGTCCTTGGGAACACTGCATCGACAGATACCCGGTCAAAATTGAT 1370
Atfael	
BnFAE1	TCTGATTCAGGTAAGTCAGAGACTCGTGTCCAAAACGGTCGGT
AtFAE1	
BnFAEl	
DHEACT	GCTCTCTTTCGTTTCTTTTATTTGTTATAATAATTTGATGGCTACGATGTTTCTCTTGT 1490
AtFAE1	
BnFAEl	TTGTTATGAATAAAGAATGCAATGGTGTTCTAGTATTTGATTGTTTTACATGTATGT
ALFAE1	
ROFAEI	

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